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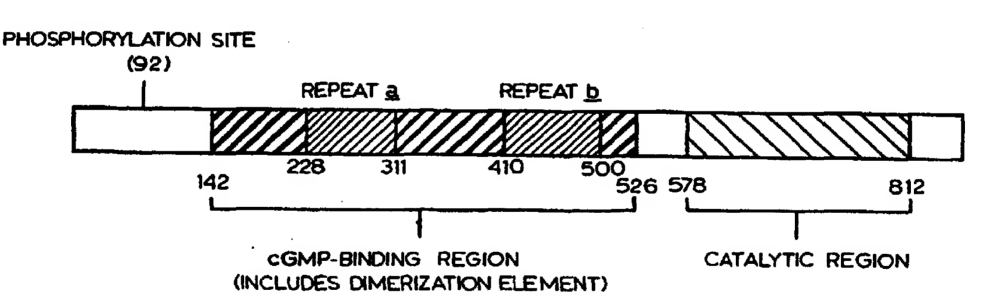
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(57) Abstract

The present invention provides novel purified and isolated nucleotide sequences encoding the cGMP-binding, cGMP-specific phosphodiesterase designated cGB-PDE. Also provided by the invention are methods and materials for the recombinant production of cGB-PDE polypeptide products and methods for identifying compounds which modulate the enzymatic activity of cGB-PDE polypeptides.

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CYCLIC GMP-BINDING, CYCLIC GMP-SPECIFIC PHOSPHODIESTERASE MATERIALS AND METHODS

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This application is a continuation-in-part of co-pending U.S. Patent Application Serial No. 08/068,051 filed May 27, 1993.

Experimental work described herein was supported in part by Research Grants GM15731, DK21723, DK40029 and GM41269 and the Medical Scientist Training Program Grant GM07347 awarded by the National Institutes of Health. The United States government has certain rights in the invention.

FIELD OF THE INVENTION

The present invention relates generally to a cyclic guanosine monophosphate-binding, cyclic guanosine monophosphate-specific phosphodiesterase designated cGB-PDE and more particularly to novel purified and isolated polynucleotides encoding cGB-PDE polypeptides, to methods and materials for recombinant production of cGB-PDE polypeptides, and to methods for identifying modulators of cGB-PDE activity.

BACKGROUND

Cyclic nucleotide phosphodiesterases (PDEs) that catalyze the hydrolysis of 3'5' cyclic nucleotides such as cyclic guanosine monophosphate (cGMP) and cyclic adenosine monophosphate (cAMP) to the corresponding nucleoside 5' monophosphates constitute a complex family of enzymes. By mediating the intracellular concentration of the cyclic nucleotides, the PDE isoenzymes function in signal transduction pathways involving cyclic nucleotide second messengers.

A variety of PDEs have been isolated from different tissue sources and many of the PDEs characterized to date exhibit differences in biological properties including physicochemical properties, substrate specificity, sensitivity to inhibitors, immunological reactivity and mode of regulation. [See Beavo et al., Cyclic Nucleotide Phosphodiesterases: Structure, Regulation and Drug Action, John Wiley & Sons, Chichester, U.K. (1990)] Comparison of the known amino acid sequences of various PDEs indicates that most PDEs are chimeric multidomain proteins that

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have distinct catalytic and regulatory domains. [See Charbonneau, pp. 267-296 in Beavo et al., supra] All mammalian PDEs characterized to date share a sequence of approximately 250 amino acid residues in length that appears to comprise the catalytic site and is located in the carboxyl terminal region of the enzyme. PDE domains that interact with allosteric or regulatory molecules are thought to be located within the amino-terminal regions of the isoenzymes. Based on their biological properties, the PDEs may be classified into six general families: the Ca²⁺/calmodulin-stimulated PDEs (Type I), the cGMP-stimulated PDEs (Type II), the cGMP-inhibited PDEs (Type III), the cAMP-specific PDEs (Type IV), the cGMP-specific phosphodiesterase cGB-PDE (Type V) which is the subject of the present invention and the cGMP-specific photoreceptor PDEs (Type VI).

The cGMP-binding PDEs (Type II, Type V and Type VI PDEs), in addition to having a homologous catalytic domain near their carboxyl terminus, have a second conserved sequence which is located closer to their amino terminus and which may comprise an allosteric cGMP-binding domain. See Charbonneau et al., Proc. Natl. Acad. Sci. USA, 87: 288-292 (1990).

The Type II cGMP-stimulated PDEs (cGs-PDEs) are widely distributed in different tissue types and are thought to exist as homodimers of 100-105 kDa subunits. The cGs-PDEs respond under physiological conditions to elevated cGMP concentrations by increasing the rate of cAMP hydrolysis. The amino acid sequence of a bovine heart cGs-PDE and a partial cDNA sequence of a bovine adrenal cortex cGS-PDE are reported in LeTrong et al., Biochemistry, 29: 10280-10288 (1990) and full length bovine adrenal and human fetal brain cGB-PDE cDNA sequences are described in Patent Cooperation Treaty International Publication No. WO 92/18541 published on October 29, 1992. The full length bovine adrenal cDNA sequence is also described in Sonnenburg et al., J. Biol. Chem., 266: 17655-17661 (1991).

The photoreceptor PDEs and the cGB-PDE have been described as cGMP-specific PDEs because they exhibit a 50-fold or greater selectivity for hydrolyzing cGMP over cAMP.

The photoreceptor PDEs are the rod outer segment PDE (ROS-PDE) and the cone PDE (COS-PDE). The holoenzyme structure of the ROS-PDE consists of two large subunits α (88 kDa) and β (84 kDa) which are both catalytically active

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and two smaller γ regulatory subunits (both 11 kDa). A soluble form of the ROS-PDE has also been identified which includes α , β , and γ subunits and a δ subunit (15 kDa) that appears to be identical to the COS-PDE 15 kDa subunit. A full-length cDNA corresponding to the bovine membrane-associated ROS-PDE α subunit is described in Ovchinnikov *et al.*, *FEBS Lett.*, 223: 169-173 (1987) and a full length cDNA corresponding to the bovine rod outer segment PDE β subunit is described in Lipkin *et al.*, *J. Biol. Chem.*, 265: 12955-12959 (1990). Ovchinnikov *et al.*, *FEBS Lett.*, 204: 169-173 (1986) presents a full-length cDNA corresponding to the bovine ROS-PDE γ subunit and the amino acid sequence of the δ subunit. Expression of the ROS-PDE has also been reported in brain in Collins *et al.*, *Genomics*, 13: 698-704 (1992). The COS-PDE is composed of two identical α' (94 kDa) subunits and three smaller subunits of 11 kDa, 13 kDa and 15 kDa. A full-length cDNA corresponding to the bovine COS-PDE α' subunit is reported in Li *et al.*, *Proc. Natl. Acad. Sci. USA*, 87: 293-297 (1990).

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cGB-PDE has been purified to homogeneity from rat [Francis et al., Methods Enzymol., 159: 722-729 (1988)] and bovine lung tissue [Thomas et al., J. Biol. Chem., 265: 14964-14970 (1990), hereinafter "Thomas I"]. The presence of this or similar enzymes has been reported in a variety of tissues and species including rat and human platelets [Hamet et al., Adv. Cyclic Nucleotide Protein Phosphorylation Res., 16: 119-136 (1984)], rat spleen [Coquil et al., Biochem. Biophys. Res. Commun., 127: 226-231 (1985)], guinea pig lung [Davis et al., J. Biol. Chem., 252: 4078-4084 (1977)], vascular smooth muscle [Coquil et al., Biochim. Biophys. Acta, 631: 148-165 (1980)], and sea urchin sperm [Francis et al., J. Biol. Chem., 255: 620-626 (1979)]. cGB-PDE may be a homodimer comprised of two 93 kDa subunits. [See Thomas I, supra] cGB-PDE has been shown to contain a single site not found in other known cGMP-binding PDEs which is phosphorylated by cGMP-dependent protein kinase (cGK) and, with a lower affinity, by cAMPdependent protein kinase (cAK). [See Thomas et al., J. Biol. Chem., 265: 14971-14978 (1990), hereinafter "Thomas II"] The primary amino acid sequence of the phosphorylation site and of the amino-terminal end of a fragment generated by chymotryptic digestion of cGB-PDE are described in Thomas II, supra, and Thomas

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I, supra, respectively. However, the majority of the amino acid sequence of cGB-PDE has not previously been described.

Various inhibitors of different types of PDEs have been described in the literature. Two inhibitors that exhibit some specificity for Type V PDEs are zaprinast and dipyridamole. See Francis et al., pp. 117-140 in Beavo et al., supra.

Elucidation of the DNA and amino acid sequences encoding the cGB-PDE and production of cGB-PDE polypeptide by recombinant methods would provide information and material to allow the identification of novel agents that selectively modulate the activity of the cGB-PDEs. The recognition that there are distinct types or families of PDE isoenzymes and that different tissues express different complements of PDEs has led to an interest in the development of PDE modulators which may have therapeutic indications for disease states that involve signal transduction pathways utilizing cyclic nucleotides as second messengers. Various selective and non-selective inhibitors of PDE activity are discussed in Murray et al., Biochem. Soc. Trans., 20(2): 460-464 (1992). Development of PDE modulators without the ability to produce a specific PDE by recombinant DNA techniques is difficult because all PDEs catalyze the same basic reaction, have overlapping substrate specificities and occur only in trace amounts. As a result, purification to homogeneity of many PDEs is a tedious and difficult process.

There thus continues to exist a need in the art for DNA and amino acid sequence information for the cGB-PDE, for methods and materials for the recombinant production of cGB-PDE polypeptides and for methods for identifying specific modulators of cGB-PDE activity.

SUMMARY OF THE INVENTION

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The present invention provides novel purified and isolated polynucleotides (e.g., DNA sequences and RNA transcripts, both sense and antisense strands, including splice variants thereof) encoding the cGMP-binding, cGMP-specific PDE designated cGB-PDE. Preferred DNA sequences of the invention include genomic and cDNA sequences as well as wholly or partially chemically synthesized DNA sequences. DNA sequences encoding cGB-PDE that are set out in SEQ ID NO: 9 or 20 and DNA sequences which hybridize thereto under stringent conditions

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or DNA sequences which would hybridize thereto but for the redundancy of the genetic code are contemplated by the invention. Also contemplated by the invention are biological replicas (i.e., copies of isolated DNA sequences made in vivo or in vitro) of DNA sequences of the invention. Autonomously replicating recombinant constructions such as plasmid and viral DNA vectors incorporating cGB-PDE sequences and especially vectors wherein DNA encoding cGB-PDE is operatively linked to an endogenous or exogenous expression control DNA sequence and a

transcriptional terminator are also provided. Specifically illustrating expression plasmids of the invention is the plasmid hogbmet 156-2 on in E. coli strain JM109

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which was deposited with the American Type Culture Collection (ATCC), 12301 Parklawn Drive, Rockville, Maryland 20852, on May 4, 1993 as Accession No. 69296.

According to another aspect of the invention, host cells including procaryotic and eucaryotic cells, are stably transformed with DNA sequences of the invention in a manner allowing the desired polypeptides to be expressed therein. Host cells expressing cGB-PDE products can serve a variety of useful purposes. Such cells constitute a valuable source of immunogen for the development of antibody substances specifically immunoreactive with cGB-PDE. Host cells of the invention are conspicuously useful in methods for the large scale production of cGB-PDE polypeptides wherein the cells are grown in a suitable culture medium and the desired polypeptide products are isolated from the cells or from the medium in which the cells are grown by, for example, immunoaffinity purification.

cGB-PDE products may be obtained as isolates from natural cell sources or may be chemically synthesized, but are preferably produced by recombinant procedures involving host cells of the invention. Use of mammalian host cells is expected to provide for such post-translational modifications (e.g., glycosylation, truncation, lipidation and tyrosine, serine or threonine phosphorylation) as may be needed to confer optimal biological activity on recombinant expression products of the invention. cGB-PDE products of the invention may be full length polypeptides, fragments or variants. Variants may comprise cGB-PDE polypeptide analogs wherein one or more of the specified (i.e., naturally encoded) amino acids is deleted or replaced or wherein one or more nonspecified amino acids are added:

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(1) without loss of one or more of the biological activities or immunological characteristics specific for cGB-PDE; or (2) with specific disablement of a particular biological activity of cGB-PDE.

Also comprehended by the present invention are antibody substances (e.g., monoclonal and polyclonal antibodies, single chain antibodies, chimeric antibodies, CDR-grafted antibodies and the like) and other binding proteins specific for cGB-PDE. Specific binding proteins can be developed using isolated or recombinant cGB-PDE or cGB-PDE variants or cells expressing such products. Binding proteins are useful, in turn, in compositions for immunization as well as for purifying cGB-PDE polypeptides and detection or quantification of cGB-PDE polypeptides in fluid and tissue samples by known immunogical procedures. They are also manifestly useful in modulating (i.e., blocking, inhibiting or stimulating) biochemical activities of cGB-PDE, especially those activities involved in signal transduction. Anti-idiotypic antibodies specific for anti-cGB-PDE antibody substances are also contemplated.

The scientific value of the information contributed through the disclosures of DNA and amino acid sequences of the present invention is manifest. As one series of examples, knowledge of the sequence of a cDNA for cGB-PDE makes possible the isolation by DNA/DNA hybridization of genomic DNA sequences encoding cGB-PDE and specifying cGB-PDE expression control regulatory sequences such as promoters, operators and the like. DNA/DNA hybridization procedures carried out with DNA sequences of the invention under stringent conditions are likewise expected to allow the isolation of DNAs encoding allelic variants of cGB-PDE, other structurally related proteins sharing one or more of the biochemical and/or immunological properties specific to cGB-PDE, and non-human species proteins homologous to cGB-PDE. Polynucleotides of the invention when suitably labelled are useful in hybridization assays to detect the capacity of cells to synthesize cGB-PDE. Polynucleotides of the invention may also be the basis for diagnostic methods useful for identifying a genetic alteration(s) in the cGB-PDE locus that underlies a disease state or states. Also made available by the invention are antisense polynucleotides relevant to regulating expression of cGB-PDE by those cells which ordinarily express the same.

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The DNA and amino acid sequence information provided by the present invention also makes possible the systematic analysis of the structure and function of cGB-PDE and definition of those molecules with which it will interact. Agents that modulate cGB-PDE activity may be identified by incubating a putative modulator with lysate from eucaryotic cells expressing recombinant cGB-PDE and determining the effect of the putative modulator on cGB-PDE phosphodiesterase activity. In a preferred embodiment the eucaryotic cell lacks endogenous cyclic nucleotide phosphodiesterase activity. Specifically illustrating such a eucaryotic cell is the yeast strain YKS45 which was deposited with the ATCC on May 19, 1993 as Accession No. 74225. The selectivity of a compound that modulates the activity of the cGB-PDE can be evaluated by comparing its activity on the cGB-PDE to its activity on other PDE isozymes. The combination of the recombinant cGB-PDE products of the invention with other recombinant PDE products in a series of independent assays provides a system for developing selective modulators of cGB-PDE.

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Selective modulators may include, for example, antibodies and other proteins or peptides which specifically bind to the cGB-PDE or cGB-PDE nucleic acid, oligonucleotides which specifically bind to the cGB-PDE or cGB-PDE nucleic acid and other non-peptide compounds (e.g., isloated or synthetic organic molecules) which specifically react with cGB-PDE or cGB-PDE nucleic acid. Mutant forms of cGB-PDE which affect the enzymatic activity or cellular localization of the wild-type cGB-PDE are also contemplated by the invention. Presently preferred targets for the development of selective modulators include, for example: (1) the regions of the cGB-PDE which contact other proteins and/or localize the cGB-PDE within a cell, (2) the regions of the cGB-PDE which bind substrate, (3) the allosteric cGMP-binding site(s) of cGB-PDE, (4) the phosphorylation site(s) of cGB-PDE and (5) the regions of the cGB-PDE which are involved in dimerization of cGB-PDE subunits. Modulators of cGB-PDE activity may be therapeutically useful in treatment of a wide range of diseases and physiological conditions.

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BRIEF DESCRIPTION OF THE DRAWINGS

Numerous other aspects and advantages of the present invention will be apparent upon consideration of the following detailed description thereof, reference being made to the drawing wherein:

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FIGURE 1A to 1C is an alignment of the conserved catalytic domains of several PDE isoenzymes wherein residues which are identical in all PDEs listed are indicated by their one letter amino acid abbreviation in the "conserved" line, residues which are identical in the cGB-PDE and photoreceptor PDEs only are indicated by a star in the "conserved" line and gaps introduced for optimum alignment are indicated by periods;

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FIGURE 2A to 2C is an alignment of the cGMP-binding domains of several PDE isoenzymes wherein residues which are identical in all PDEs listed are indicated by their one letter amino acid abbreviation in the "conserved" line and gaps introduced for optimum alignment are indicated by periods;

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FIGURE 3 is an alignment of internally homologous repeats from several PDE isoenzymes wherein residues identical in each repeat A and B from all cGMP-binding PDEs listed are indicated by their one letter amino acid abbreviation in the "conserved" line and stars in the "conserved" line represent positions in which all residues are chemically conserved;

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FIGURE 4 schematically depicts the domain organization of cGB-PDE; FIGURE 5 is a bar graph representing the results of experiments in which extracts of COS cells transfected with bovine cGB-PDE sequences or extracts of untransfected COS cells were assayed for phosphodiesterase activity using either 20 μ M cGMP or 20 μ M cAMP as the substrate;

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FIGURE 6 is a graph depicting results of assays of extracts from cells transfected with bovine cGB-PDE sequences for cGMP phosphodiesterase activity in the presence of a series of concentrations of phosphodiesterase inhibitors including dypyridamole (closed squares), zaprinast (closed circles), methoxymethylxanthine (closed triangles) and rolipram (open circles);

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FIGURE 7 is a bar graph presenting results of experiments in which cell extracts from COS cells transfected with bovine cGB-PDE sequences or control

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untransfected COS cells were assayed for [3H]cGMP-binding activity in the absence (-) or presence (+) of 0.2 mM IBMX; and

FIGURE 8 is a graph of the results of assays in which extracts from cells transfected with bovine cGB-PDE sequences were assayed for [3H]cGMP-binding activity in the presence of excess unlabelled cAMP (open circles) or cGMP (closed circles) at the concentrations indicated.

DETAILED DESCRIPTION

The following examples illustrate the invention. Example 1 describes the isolation of a bovine cGB-PDE cDNA fragment by PCR and subsequent isolation of a full length cGB-PDE cDNA using the PCR fragment as a probe. Example 2 presents an analysis of the relationship of the bovine cGB-PDE amino acid sequence to sequences reported for various other PDEs. Northern blot analysis of cGB-PDE mRNA in various bovine tissues is presented in Example 3. Expression of the bovine cGB-PDE cDNA in COS cells is described in Example 4. Example 5 presents results of assays of the cGB-PDE COS cell expression product for phosphodiesterase activity, cGMP-binding activity and Zn²⁺ hydrolase activity. Example 6 describes the isolation of human cDNAs homologous to the bovine cGB-PDE cDNA. expression of a human cGB-PDE cDNA in yeast cells is presented in Example 7. RNase protection assays to detect cGB-PDE in human tissues are described in Example 8. Example 9 describes the bacterial expression of human cGB-PDE cDNA and the development of antibodies reactive with the bacterial cGB-PDE expression product. Example 10 describes cGB-PDE analogs and fragments. The generation of monoclonal antibodies that recognize cGB-PDE is described in Example 11. Example 12 relates to utilizing recombinant cGB-PDE products of the invention to develop agents that selectively modulate the biological activities of cGB-PDE.

Example 1

The polymerase chain reaction (PCR) was utilized to isolate a cDNA fragment encoding a portion of cGB-PDE from bovine lung first strand cDNA. Fully degenerate sense and antisense PCR primers were designed based on the partial cGB-

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PDE amino acid sequence described in Thomas I, supra, and novel partial amino acid sequence information.

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A. Purification of cGB-PDE Protein

cGB-PDE was purified as described in Thomas I, supra, or by a modification of that method as described below.

Fresh bovine lungs (5-10 kg) were obtained from a slaughterhouse and immediately placed on ice. The tissue was ground and combined with cold PEM buffer (20mM sodium phosphate, pH 6.8, containing 2mM EDTA and 25mM β mercaptoethanol). After homogenization and centrifugation, the resulting supernatant was incubated with 4-7 liters of DEAE-cellulose (Whatman, UK) for 3-4 hours. The DEAE slurry was then filtered under vacuum and rinsed with multiple volumes of cold PEM. The resin was poured into a glass column and washed with three to four volumes of PEM. The protein was eluted with 100mM NaCl in PEM and twelve 1liter fractions were collected. Fractions were assayed for IBMX-stimulated cGMP binding and cGMP phosphodiesterase activities by standard procedures described in Thomas et al., supra. Appropriate fractions were pooled, diluted 2-fold with cold, deionized water and subjected to Blue Sepharose® CL-6B (Pharmacia LKB) Biotechnology Inc., Piscataway, NJ) chromatography. Zinc chelate affinity adsorbent chromatography was then performed using either an agarose or Sepharose-based gel matrix. The resulting protein pool from the zinc chelation step treated as described in the Thomas I, supra, or was subjected to a modified purification procedure.

As decribed in Thomas I, supra, the protein pool was applied in multiple loads to an HPLC Bio-Sil TSK-545 DEAE column (150 x 21.5 mm) (BioRad Laboratories, Hercules, CA) equilibrated in PEM at 4°C. After an equilibration period, a 120-ml wash of 50mM NaCl in PEM was followed by a 120-ml linear gradient (50-200mM NaCl in PEM) elution at a flow rate of 2 ml/minute. Appropriate fractions were pooled and concentrated in dialysis tubing against Sephadex G-200 (Boehringer Mannheim Biochemicals, UK) to a final volume of 1.5 ml. The concentrated cGB-PDE pool was applied to an HPLC gel filtration column (Bio-Sil TSK-250, 500 x 21.5 mm) equilibrated in 100mM sodium phosphate, pH 6.8, 2mM EDTA, 25mM β -mercaptoethanol and eluted with a flow rate of 2 ml/minute at 4°C.

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If the modified, less cumbersome procedure was performed, the protein pool was dialyzed against PEM for 2 hours and loaded onto a 10 ml preparative DEAE Sephacel column (Pharmacia) equilibrated in PEM buffer. The protein was eluted batchwise with 0.5M NaCl in PEM, resulting in an approximately 10-15 fold concentration of protein. The concentrated protein sample was loaded onto an 800 ml (2.5 cm x 154 cm) Sephacryl S400 gel filtration column (Boehringer) equilibrated in 0.1M NaCl in PEM, and eluted at a flow rate of 1.7 ml/minute.

The purity of the protein was assessed by Coomassie staining after sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE). Approximately 0.5-3.0 mg of pure cGB-PDE were obtained per 10 kg bovine lung.

Rabbit polyclonal antibodies specific for the purified bovine cGB-PDE were generated by standard procedures.

B. Amino Acid Sequencing of cGB-PDE

cGB-PDE phosphorylated with [32 P]ATP and was then digested with protease to yield 32 P-labelled phosphopeptides. Approximately 100 μ g of purified cGB-PDE was phosphorylated in a reaction mixture containing 9mM MgCl₂, 9 μ M [32 P]ATP, 10 μ M cGMP, and 4.2 μ g purified bovine catalytic subunit of cAMP-dependent protein kinase (cAK) in a final volume of 900 μ l. Catalytic subunit of cAK was prepared according to the method of Flockhart *et al.*, pp. 209-215 in Marangos et al., *Brain Receptor Methodologies*, *Part A*, Academic Press, Orlando, Florida (1984). The reaction was incubated for 30 minutes at 30°C, and stopped by addition of 60 μ l of 200mM EDTA.

To obtain a first peptide sequence from cGB-PDE, 3.7 μ l of a 1 mg/ml solution of a α -chymotrypsin in KPE buffer (10mM potassium phosphate, pH 6.8, with 2mM EDTA) was added to 100 μ g purified, phosphorylated cGB-PDE and the mixture was incubated for 30 minutes at 30°C. Proteolysis was stopped by addition of 50 μ l of 10% SDS and 25 μ l of β -mercaptoethanol. The sample was boiled until the volume was reduced to less than 400 μ l, and was loaded onto an 8% preparative SDS-polyacrylamide gel and subjected to electrophoresis at 50mAmps. The separated digestion products were electroblotted onto Immobilon polyvinylidene difluoride (Millipore, Bedford, MA), according to the method of Matsudaira, *J. Biol. Chem*, 262: 10035-10038 (1987). Transferred protein was identified by Coomassie Blue

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staining, and a 50 kDa band was excised from the membrane for automated gas-phase amino acid sequencing. The sequence of the peptide obtained by the α -chymotryptic digestion procedure is set out below as SEQ ID NO: 1.

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SEQ ID NO: 1

REXDANRINYMYAQYVKNTM

A second sequence was obtained from a cGB-PDE peptide fragment generated by V8 proteolysis. Approximately 200 μ g of purified cGB-PDE was added to 10mM MgCl₂, 10μ M [32 P]ATP, 100μ M cGMP, and $1~\mu$ g/ml purified catalytic subunit of cAK in a final volume of 1.4 ml. The reaction was incubated for 30 minutes at 30°C, and was terminated by the addition of 160 μ l of 0.2M EDTA. Next, 9 μ l of 1 mg/ml *Staphylococcal aureus* V8 protease (International Chemical Nuclear Biomedicals, Costa Mesa, CA) diluted in KPE was added, followed by a 15 minute incubation at 30°C. Proteolysis was stopped by addition of 88 μ l of 10% SDS and 45 μ l β -mercaptoethanol. The digestion products were separated by electrophoresis on a preparative 10% SDS-polyacrylamide gel run at 25 mAmps for 4.5 hours. Proteins were electroblotted and stained as described above. A 28 kDa protein band was excised from the membrane and subjected to automated gas-phase amino acid sequencing. The sequence obtained is set out below as SEQ ID NO: 2.

SEQ ID NO: 2

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C. PCR Amplification of Bovine cDNA

The partial amino acid sequences utilized to design primers (SEQ ID NO: 3, below, and amino acids 9-20 of SEQ ID NO: 1) and the sequences of the corresponding PCR primers (in IUPAC nomenclature) are set below wherein SEQ ID NO: 3 is the sequence reported in Thomas I, supra.

SEQ ID NO: 3

F D N D E G E Q

- 5' TTY GAY AAY GAY GAR GGN GAR CA 3' (SEQ ID NO: 4)
- 3' AAR CTR TTR CTR CTY CCN CTY GT 5' (SEQ ID NO: 5)

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SEQ ID NO: 1, Amino acids 9-20

NYMYAQYVKNTM

- 5' AAY TAY ATG TAY GCN CAR TAY GT 3' (SEQ ID NO: 6)
- 3' TTR ATR TAC ATR CGN GTY ATR CA 5' (SEQ ID NO: 7)
- 3' TTR ATR TAC ATR CGN GTY ATR CAN TTY TTR TGN TAC 5' (SEQ ID NO: 8)

The sense and antisense primers, synthesized using an Applied Biosystems Model 380A DNA Synthesizer (Foster City, CA), were used in all possible combinations to amplify cGB-PDE-specific sequences from bovine lung first strand cDNA as described below.

After ethanol precipitation, pairs of oligonucleotides were combined (SEQ ID NO: 4 or 5 combined with SEQ ID NOs: 6, 7 or 8) at 400nM each in a PCR reaction. The reaction was run using 50 ng first strand bovine lung cDNA (generated using AMV reverse transcriptase and random primers on oligo dT selected bovine lung mRNA), 200 µM dNTPs, and 2 units of Taq polymerase. The initial denaturation step was carried out at 94°C for 5 minutes, followed by 30 cycles of a 1 minute denaturation step at 94°C, a two minute annealing step at 50°C, and a 2 minute extension step at 72°C. PCR was performed using a Hybaid Thermal Reactor (ENK Scientific Products, Saratoga, CA) and products were separated by gel electrophoresis on a 1% low melting point agarose gel run in 40mM Tris-acetate, 2mM EDTA. A weak band of about 800-840 bp was seen with the primers set out in SEQ ID NOs: 4 and 7 and with primers set out in SEQ ID NOs: 4 and 8. None of the other primer pairs yielded visible bands. The PCR product generated by amplification with the primers set out in SEQ ID NOs: 4 and 7 was isolated using the Gene Clean® (Bio101, La Jolla, CA) DNA purification kit according to the manufacturer's protocol. The PCR product (20 ng) was ligated into 200 ng of linearized pBluescript KS(+) (Stratagene, La Jolla, CA), and the resulting plasmid construct was used to transform E. coli XL1 Blue cells (Stratagene Cloning Systems, La Jolla, CA). Putative transformation positives were screened by sequencing. The sequences obtained were not homologous to any known PDE sequence or to the known partial cGB-PDE sequences.

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PCR was performed again on bovine lung first strand cDNA using the primers set out in SEQ ID NOs: 4 and 7. A clone containing a 0.8 Kb insert with a single large open reading frame was identified. The open reading frame encoded a polypeptide that included the amino acids KNTM (amino acids 17-20 of SEQ ID NO: 1 which were not utilized to design the primer sequence which is set out in SEQ ID NO: 7) and that possessed a high degree of homology to the deduced amino acid sequences of the cGs-, ROS- and COS-PDEs. The clone identified corresponds to nucleotides 489-1312 of SEQ ID NO: 9.

D. Construction and Hybridization Screening

10 of a Bovine cDNA Library

In order to obtain a cDNA encoding a full-length cGB-PDE, a bovine lung cDNA library was screened using the ³²P-labelled PCR-generated cDNA insert as a probe.

Polyadenylated RNA was prepared from bovine lung as described Sonnenburg et al., J. Biol. Chem., 266: 17655-17661 (1991). First strand cDNA was synthesized using AMV reverse transcriptase (Life Sciences, St. Petersburg, FL) with random hexanucleotide primers as described in Ausubel et al., Current Protocols in Molecular Biology, John Wiley & Sons, New York (1987). Second strand cDNA was synthesized using E. coli DNA polymerase I in the presence of E. coli DNA ligase and E. coli RNAse H. Selection of cDNAs larger than 500 bp was performed by Sepharose® CL-4B (Millipore) chromatography. EcoRI adaptors (Promega, Madison, WI) were ligated to the cDNA using T4 DNA ligase. Following heat inactivation of the ligase, the cDNA was phosphorylated using T4 polynucleotide kinase. Unligated adaptors were removed by Sepharose® CL-4B chromatography (Pharmacia, Piscataway, NJ). The cDNA was ligated into EcoRI-digested, dephosphorylated lambda Zap®II arms (Stratagene) and packaged with Gigapack® Gold (Stratagene) extracts according to the manufacturer's protocol. The titer of the unamplified library was 9.9 x 10⁵ with 18% nonrecombinants. The library was amplified by plating 50,000 plaque forming units (pfu) on to twenty 150 mm plates, resulting in a final titer of 5.95 x 10⁶ pfu/ml with 21% nonrecombinants.

The library was plated on twenty-four 150 mm plates at 50,000 pfu/plate, and screened with the ³²P-labelled cDNA clone. The probe was prepared

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using the method of Feinberg *et al.*, *Anal. Biochem.*, 137: 266-267 (1984), and the 32 P-labelled DNA was purified using Elutip-D® columns (Schleicher and Schuell Inc., Keene, NH) using the manufacturer's protocol. Plaque-lifts were performed using 15 cm nitrocellulose filters. Following denaturation and neutralization, DNA was fixed onto the filters by baking at 80°C for 2 hours. Hybridization was carried out at 42°C overnight in a solution containing 50% formamide, 5X SSC (0.75M NaCl, 0.75M sodium citrate, pH 7), 25mM sodium phosphate (pH 7.0), 2X Denhardt's solution, 10% dextran sulfate, 90 μ g/ml yeast tRNA, and approximately 106 cpm/ml 32 P-labelled probe (5x108 cpm/ μ g). The filters were washed twice in 0.1X SSC, 0.1% SDS at room temperature for 15 minutes per wash, followed by a single 20 minute wash in 0.1X SSC, 1% SDS at 45°C. The filters were then exposed to X-ray film at -70°C for several days.

Plaques that hybridized with the labelled probe were purified by several rounds of replating and rescreening. Insert cDNAs were subcloned into the pBluescript SK(-) vector (Stratagene) by the *in vivo* excision method described by the manufacturer's protocol. Southern blots were performed in order to verify that the rescued cDNA hybridized to the PCR probe. Putative cGB-PDE cDNAs were sequenced using Sequenase® Version 2.0 (United States Biochemical Corporation, Cleveland, Ohio) or TaqTrack® kits (Promega).

Three distinct cDNA clones designated cGB-2, cGB-8 and cGB-10 were isolated. The DNA and deduced amino acid sequences of clone cGB-8 are set out in SEQ ID NOs: 9 and 10. The DNA sequence downstream of nucleotide 2686 may represent a cloning artifact. The DNA sequence of cGB-10 is identical to the sequence of cGB-8 with the exception of one nucleotide. The DNA sequence of clone cGB-2 diverges from that of clone cGB-8 5' to nucleotide 219 of clone cgb-8 (see SEQ ID NO: 9) and could encode a protein with a different amino terminus.

The cGB-8 cDNA clone is 4474 bp in length and contains a large open reading frame of 2625 bp. The triplet ATG at position 99-101 in the nucleotide sequence is predicted to be the translation initiation site of the cGB-PDE gene because it is preceded by an in-frame stop codon and the surrounding bases are compatible with the Kozak consensus initiation site for eucaryotic mRNAs. The stop codon TAG is located at positions 2724-2726, and is followed by 1748 bp of 3' untranslated

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sequence. The sequence of cGB-8 does not contain a transcription termination consensus sequence, therefore the clone may not represent the entire 3' untranslated region of the corresponding mRNA.

The open reading frame of the cGB-8 cDNA encodes an 875 amino acid polypeptide with a calculated molecular mass of 99.5 kD. This calculated molecular mass is only slightly larger than the reported molecular mass of purified cGB-PDE, estimated by SDS-PAGE analysis to be approximately 93 kDa. The deduced amino acid sequence of cGB-8 corresponded exactly to all peptide sequences obtained from purified bovine lung cGB-PDE providing strong evidence that cGB-8 encodes cGB-PDE.

Example 2

A search of the SWISS-PROT and GEnEmbl data banks (Release of February, 1992) conducted using the FASTA program supplied with the Genetics Computer Group (GCG) Software Package (Madison, Wisconsin) revealed that only DNA and amino acid sequences reported for other PDEs displayed significant similarity to the DNA and deduced amino acid of clone cGB-8.

Pairwise comparisons of the cGB-PDE deduced amino acid sequence with the sequences of eight other PDEs were conducted using the ALIGN [Dayhoff et al., Methods Enzymol., 92: 524-545 (1983)] and BESTFIT [Wilbur et al., Proc. Natl. Acad. Sci. USA, 80: 726-730 (1983)] programs. Like all mammalian phosphodiesterases sequenced to date, cGB-PDE contains a conserved catalytic domain sequence of approximately 250 amino acids in the carboxyl-terminal half of the protein that is thought to be essential for catalytic activity. This segment comprises amino acids 578-812 of SEQ ID NO: 9 and exhibits sequence conservation with the corresponding regions of other PDEs. Table 1 below sets out the specific identity values obtained in pairwise comparisons of other PDEs with amino acids 578-812 of cGB-PDE, wherein "ratdunce" is the rat cAMP-specific PDE; "61 kCaM" is the bovine 61 kDa calcium/calmodulin-dependent PDE; "drosdunce" is the drosophila cAMP-specific dunce PDE; "ROS- α " is the bovine ROS-PDE α -subunit; "ROS- β " is the

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bovine ROS-PDE β -subunit; "COS- α '" is the bovine COS-PDE α ' subunit; and "cGs" is the bovine cGs-PDE (612-844).

Table 1

	Phosphodiesterase	Catalytic Domain Residues	% Identity
5	Ratdunce	77-316	31
	61 kCaM	193-422	29
	63 kcam	195-424	29
	drosdunce	1-239	28
	ROS-α	535-778	45
10	ROS-β	533-776	46
	COS-α'	533-776	48
	cGs	612-844	40

Multiple sequence alignments were performed using the Progressive Alignment Algorithm [Feng et al., Methods Enzymol., 183: 375-387 (1990)] implemented in the PILEUP program (GCG Software). FIGURE 1A to 1C shows a multiple sequence alignment of the proposed catalytic domain of cGB-PDE with the all the corresponding regions of the PDEs of Table 1. Twenty-eight residues (see residues indicated by one letter amino acid abbreviations in the "conserved" line on FIGURE 1A to 1C) are invariant among the isoenzymes including several conserved histidine residues predicted to play a functional role in catalysis. See Charbonneau et al., Proc. Natl. Acad. Sci. USA, supra. The catalytic domain of cGB-PDE more closely resembles the catalytic domains of the ROS-PDEs and COS-PDEs than the corresponding regions of other PDE isoenzymes. There are several conserved regions among the photoreceptor PDEs and cGB-PDE that are not shared by other PDEs. Amino acid positions in these regions that are invariant in the photoreceptor PDE and cGB-PDE sequences are indicated by stars in the "conserved" line of FIGURE 1A to 1C. Regions of homology among cGB-PDE and the ROS- and COS-PDEs may serve important roles in conferring specificity for cGMP hydrolysis relative to cAMP hydrolysis or for sensitivity to specific pharmacological agents.

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Sequence similarity between cGB-PDE, cGs-PDE and the photoreceptor PDEs, is not limited to the conserved catalytic domain but also includes the noncatalytic cGMP binding domain in the amino-terminal half of the protein. Optimization of the alignment between cGB-PDE, cGs-PDE and the photoreceptor PDEs indicates that an amino-terminal conserved segment may exist including amino acids 142-526 of SEQ ID NO: 9. Pairwise analysis of the sequence of the proposed cGMP-binding domain of cGB-PDE with the corresponding regions of the photoreceptor PDEs and cGs-PDE revealed 26-28% sequence identity. Multiple sequence alignment of the proposed cGMP-binding domains with the cGMP-binding PDEs is shown in FIGURE 2A to 2C wherein abbreviations are the same as indicated for Table 1. Thirty-eight positions in this non-catalytic domain appear to be invariant among all cGMP-binding PDEs (see positions indicated by one letter amino acid abbreviations in the "conserved" line of FIGURE 2A to 2C).

The cGMP-binding domain of the cGMP-binding PDEs contains internally homologous repeats which may form two similar but distinct inter- or intrasubunit cGMP-binding sites. FIGURE 3 shows a multiple sequence alignment of the repeats a (corresponding to amino acids 228-311 of cGB-PDE) and b (corresponding to amino acids 410-500 of cGB-PDE) of the cGMP-binding PDEs. Seven residues are invariant in each A and B regions (see residues indicated by one letter amino acid abbreviations in the "conserved" line of FIGURE 3). Residues that are chemically conserved in the A and B regions are indicated by stars in the "conserved" line of FIGURE 3. cGMP analog studies of cGB-PDE support the existence of a hydrogen bond between the cyclic nucleotide binding site on cGB-PDE and the 2'OH of cGMP.

Three regions of cGB-PDE have no significant sequence similarity to other PDE isoenzymes. These regions include the sequence flanking the carboxylterminal end of the catalytic domain (amino acids 812-875), the sequence separating the cGMP-binding and catalytic domains (amino acids 527-577) and the aminoterminal sequence spanning amino acids 1-141. The site (the serine at position 92 of SEQ ID NO: 10) of phosphorylation of cGB-PDE by cGK is located in this aminoterminal region of sequence. Binding of cGMP to the allosteric site on cGB-PDE is required for its phosphorylation.

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A proposed domain structure of cGB-PDE based on the foregoing comparisons with other PDE isoenzymes is presented in FIGURE 4. This domain structure is supported by the biochemical studies of cGB-PDE purified from bovine lung.

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Example 3

The presence of cGB-PDE mRNA in various bovine tissues was examined by Northern blot hybridization.

Polyadenylated RNA was purified from total RNA preparations using the Poly(A) Quick® mRNA purification kit (Stratagene) according to the manufacturer's protocol. RNA samples (5 μ g) were loaded onto a 1.2% agarose, 6.7% formaldehyde gel. Electrophoresis and RNA transfer were performed as previously described in Sonnenburg et al., supra. Prehybridization of the RNA blot was carried out for 4 hours at 45°C in a solution containing 50% formamide, 5X SSC, 25mM sodium phosphate, pH 7, 2X Denhardt's solution, 10% dextran sulfate, and 0.1 mg/ml yeast tRNA. A random hexanucleotide-primer-labelled probe (5 X 10^8 cpm/ μ g) was prepared as described in Feinberg et al., supra, using the 4.7 kb cGB-8 cDNA clone of Example 2 excised by digestion with AccI and SacII. The probe was heat denatured and injected into a blotting bag (6 X 10⁵ cpm/ml) following prehybridization. The Northern blot was hybridized overnight at 45°C, followed by one 15 minute wash with 2X SSC, 0.1% SDS at room temperature, and three 20 minute washes with 0.1X SSC, 0.1% SDS at 45°C. The blot was exposed to X-ray film for 24 hours at -70°C. The size of the RNA that hybridized with the cGB-PDE probe was estimated using a 0.24-9.5 kb RNA ladder that was stained with ethidium bromide and visualized with UV light.

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The ³²P-labelled cGB-PDE cDNA hybridized to a single 6.8 kb bovine lung RNA species. A mRNA band of the identical size was also detected in polyadenylated RNA isolated from bovine trachea, aorta, kidney and spleen.

Example 4

The cGB-PDE cDNA in clone cGB-8 of Example 2 was expressed in COS-7 cells (ATCC CRL1651).

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A portion of the cGB-8 cDNA was isolated following digestion with the restriction enzyme XbaI. XbaI cut at a position in the pBluescript polylinker sequence located 30 bp upstream of the 5' end of the cGB-8 insert and at position 3359 within the cGB-8 insert. The resulting 3389 bp fragment, which contains the entire coding region of cGB-8, was then ligated into the unique XbaI cloning site of the expression vector pCDM8 (Invitrogen, San Diego, CA). The pCDM8 plasmid is a 4.5 kb eucaryotic expression vector containing a cytomegalovirus promoter and enhancer, an SV40-derived origin of replication, a polyadenylation signal, a procaryotic origin of replication (derived from pBR322) and a procaryotic genetic marker (supF). E. coli MC1061/P3 cells (Invitrogen) were transformed with the resulting ligation products, and transformation positive colonies were screened for proper orientation of the cGB-8 insert using PCR and restriction enzyme analysis. The resulting expression construct containing the cGB-8 insert in the proper orientation is referred to as pCDM8-cGB-PDE.

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The pCDM8-cGB-PDE DNA was purified from large-scale plasmid preparations using Qiagen pack-500 columns (Chatsworth, CA) according to the manufacturer's protocol. COS-7 cells were cultured in Dulbecco's modified Eagle's medium (DMEM) containing 10% fetal bovine serum, 50 μ g/ml penicillin and 50 μg/ml streptomycin at 37°C in a humidified 5% CO₂ atmosphere. Approximately 24 hours prior to transfection, confluent 100 mm dishes of cells were replated at onefourth or one-fifth the original density. In a typical transfection experiment, cells were washed with buffer containing 137mM NaCl, 2.7mM KCl, 1.1mM potassium phosphate, and 8.1mM sodium phosphate, pH 7.2 (PBS). Then 4-5 ml of DMEM containing 10% NuSerum (Collaborative Biomedical Products, Bedford, MA) was added to each plate. Transfection with 10 μ g pCDM8-cGB-PDE DNA or pCDM8 vector DNA mixed with 400 μg DEAE-dextran (Pharmacia) in 60 μl TBS [Trisbuffered saline: 25mM Tris-HCl (pH 7.4), 137mM NaCl, 5mM KCl, 0.6mM Na₂HPO₄, 0.7mM CaCl₂, and 0.5mM MgCl₂] was carried out by dropwise addition of the mixture to each plate. The cells were incubated at 37°C, 5% CO₂ for 4 hours, and then treated with 10% dimethyl sulfoxide in PBS for 1 minute. After 2 minutes, the dimethyl sulfoxide was removed, the cells were washed with PBS and incubated in complete medium. After 48 hours, cells were suspended in 0.5-1 ml of cold

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homogenization buffer [40mM Tris-HCl (pH 7.5), 15mM benzamidine, 15mM β -mercaptoethanol, 0.7 μ g/ml pepstatin A, 0.5 μ g/ml leupeptin, and 5 μ M EDTA] per plate of cells, and disrupted using a Dounce homogenizer. The resulting whole-cell extracts were assayed for phosphodiesterase activity, cGMP-binding activity, and total protein concentration as described below in Example 5.

Example 5

Phosphodiesterase activity in extracts of the transfected COS cells of Example 4 or in extracts of mock transfected COS cells was measured using a modification of the assay procedure described for the cGs-PDE in Martins *et al.*, *J. Biol. Chem.*, 257: 1973-1979 (1982). Cells were harvested and extracts prepared 48 hours after transfection. Incubation mixtures contained 40mM MOPS buffer (pH 7), 0.8mM EDTA, 15mM magnesium acetate, 2 mg/ml bovine serum albumin, 20μ M [3 H]cGMP or [3 H]cAMP (100,000-200,000 cpm/assay) and COS-7 cell extract in a total volume of 250 μ l. The reaction mixture was incubated for 10 minutes at 30 °C, and then stopped by boiling. Next, 10μ l of 10mg/ml *Crotalus atrox* venom (Sigma) was added followed by a 10 minute incubation at 30 °C. Nucleoside products were separated from unreacted nucleotides as described in Martins *et al.*, *supra*. In all studies, less than 15% of the total [3 H]cyclic nucleotide was hydrolyzed during the reaction.

The results of the assays are presented in FIGURE 5 wherein the results shown are averages of three separate transfections. Transfection of COS-7 cells with pCDM8-cGB-PDE DNA resulted in the expression of approximately 15-fold higher levels of cGMP phosphodiesterase activity than in mock-transfected cells or in cells transfected with pCDM8 vector alone. No increase in cAMP phosphodiesterase activity over mock or vector-only transfected cells was detected in extracts from cells transfected with pCDM8-cGB-PDE DNA. These results confirm that the cGB-PDE bovine cDNA encodes a cGMP-specific phosphodiesterase.

Extracts from the transfected COS cells of Example 4 were also assayed for cGMP PDE activity in the presence of a series of concentrations of the PDE inhibitors zaprinast, dipyridamole (Sigma), isobutyl-1-methyl-8-methoxymethylxanthine (MeOxMeMIX) and rolipram.

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The results of the assays are presented in FIGURE 6 wherein PDE activity in the absence of inhibitor is taken as 100% and each data point represents the average of two separate determinations. The relative potencies of PDE inhibitors for inhibition of cGMP hydrolysis by the expressed cGB-BPDE cDNA protein product were identical to those relative potencies reported for native cGB-PDE purified from bovine lung (Thomas I, supra). IC₅₀ values calculated from the curves in FIGURE 6 are as follows: zaprinast (closed circles), 2 μ M; dipyridamole (closed squares), 3.5 μ M; MeOxMeMIX (closed triangles), 30 μ M; and rolipram (open circles), $>300 \mu M$. The IC₅₀ value of zaprinast, a relatively specific inhibitor of cGMP-specific phosphodiesterases, was at least two orders of magnitude lower than that reported for inhibition of phosphodiesterase activity of the cGs-PDE or of the cGMP-inhibited phosphodiesterase (cGi-PDEs) (Reeves et al., pp. 300-316 in Beavo et al., supra). Dipyrimadole, an effective inhibitor of selected cAMP- and cGMPspecific phosphodiesterases, was also a potent inhibitor of the expressed cGB-PDE. The relatively selective inhibitor of calcium/calmodulin-stimulated phosphodiesterase (CaM-PDEs), MeOxMeMIX, was approximately 10-fold less potent than zaprinast and dipyridamole, in agreement with results using cGB-PDE activity purified from bovine lung. Rolipram, a potent inhibitor of low K_m cAMP phosphodisterases, was a poor inhibitor of expressed cGB-PDE cDNA protein product. These results show that the cGB-PDE cDNA encodes a phosphodiesterase that possesses catalytic activity characteristic of cGB-PDE isolated from bovine tissue, thus verifying the identity of the cGB-8 cDNA clone as a cGB-PDE.

It is of interest to note that although the relative potencies of the PDE inhibitors for inhibition of cGMP hydrolysis were identical for the recombinant and bovine isolate cGB-PDE, the absolute IC₅₀ values for all inhibitors tested were 2-7 fold higher for the recombinant cGB-PDE. This difference could not be attributed to the effects of any factors present in COS-7 cell extracts on cGMP hydrolytic activity, since cGB-PDE isolated from bovine tissue exhibited identical kinetics of inhibition as a pure enzyme, or when added back to extracts of mock-transfected COS-7 cells. This apparent difference in pharmacological sensitivity may be due to a subtle difference in the structure of the recombinant cGB-PDE cDNA protein product and bovine lung cGB-PDE, such as a difference in post-translational

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modification at or near the catalytic site. Alternatively, this difference may be due to an alteration of the catalytic activity of bovine lung cGB-PDE over several purification steps.

Cell extracts were assayed for [3H]cGMP-binding activity in the absence or presence of 0.2mM 3-isobutyl-1-methylaxanthine (IBMX) (Sigma), a competitive inhibitor of cGMP hydrolysis. The cGMP binding assay, modified from the assay described in Thomas I, supra, was conducted in a total volume of 80 μ l. Sixty μ l of cell extract was combined with 20 μ l of a binding cocktail such that the final concentration of components of the mixture were $1\mu M$ [3H]cGMP, $5\mu M$ cAMP, and 10µM 8-bromo-cGMP. The cAMP and 8-bromo-cGMP were added to block [3H]cGMP binding to cAK and cGK, respectively. Assays were carried out in the absence and presence of 0.2mM IBMX. The reaction was initiated by the addition of the cell extract, and was incubated for 60 minutes at 0°C. Filtration of the reaction mixtures was carried out as described in Thomas I, supra. Blanks were determined by parallel incubations with homogenization buffer replacing cell extracts, or with a 100-fold excess of unlabelled cGMP. Similar results were obtained with both methods. Total protein concentration of the cell extracts was determined by the method of Bradford, Anal. Biochem., 72:248-254 (1976) using bovine serum albumin as the standard.

Results of the assay are set out in FIGURE 7. When measured at 1μM [³H]cGMP in the presence of 0.2mM IBMX, extracts from COS-7 cells transfected with pCDM8-cGB-PDE exhibited 8-fold higher cGMP-binding activity than extracts from mock-transfected cells. No IBMX stimulation of background cGMP binding was observed suggesting that little or no endogenous cGB-PDE was present in the COS-7 cell extracts. In extracts of pCDM8-cGB-PDE transfected cells cGMP-specific activity was stimulated approximately 1.8-fold by the addition of 0.2mM IBMX. The ability of IBMX to stimulate cGMP binding 2-5 fold is a distinctive property of the cGMP-binding phosphodisterases.

Cell extracts were assayed as described above for [³H]cGMP-binding activity (wherein concentration of [³H]cGMP was 2.5 μ M) in the presence of excess unlabelled cAMP or cGMP. Results are presented in FIGURE 8 wherein cGMP binding in the absence of unlabelled competitor was taken as 100% and each data

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point represents the average of three separate determinations. The binding activity of the protein product encoded by the cGB-PDE cDNA was specific for cGMP relative to cAMP. Less than 10-fold higher concentrations of unlabelled cGMP were required to inhibit [³H]cGMP binding activity by 50% whereas approximately 100-fold higher concentrations of cAMP were required for the same degree of inhibition.

The results presented in this example show that the cGB-PDE cDNA encodes a phosphodiesterase which possesses biochemical activities characteristic of native cGB-PDE.

The catalytic domains of mammalian PDEs and a Drosophila PDE contain two tandem conserved sequences (HX₃HX₂₄₋₂₆E) that are typical Zn²⁺-binding motifs in Zn2+ hydrolases such as thermolysin [Vallee and Auld, Biochem., 29: 5647-5659 (1990)]. cGB-PDE binds Zn²⁺ in the presence of large excesses of Mg²⁺, Mn²⁺, Fe²⁺, Fe³⁺, Ca²⁺ or Cd²⁺. In the absence of added metal, cGB-PDE has a PDE activity that is approximately 20% of the maximum activity that occurs in the presence of 40 mM Mg²⁺, and this basal activity is inhibited by 1,10-phenanthroline or EDTA. This suggests that a trace metal(s) accounts for the basal PDE activity despite exhaustive treatments to remove metals. PDE activity is stimulated by addition of Zn^{2+} (0.02-1 μ M) or Co^{2+} (1-20 μ M), but not by Fe^{2+} , Fe^{3+} , $C\alpha^{2+}$, Cd^{2+} , or Cu²⁺. Zn²⁺ increases the basal PDE activity up to 70% of the maximum stimulation produced by 40mM Mg²⁺. The stimulatory effect of Zn²⁺ in these assays may be compromised by an inhibitory effect that is caused by Zn²⁺ concentrations > 1 μ M. The Zn²⁺-supported PDE activity and Zn²⁺ binding by cGB-PDE occur at similar concentrations of Zn²⁺. cGB-PDE thus appears to be a Zn²⁺ hydrolase and Zn²⁺ appears to play a critical role in the activity of the enzyme. See, Colbran et al., The FASEB J., 8: Abstract 2148 (March 15, 1994).

Example 6

Several human cDNA clones, homologous to the bovine cDNA clone encoding cGB-PDE, were isolated by hybridization under stringent conditions using a nucleic acid probe corresponding to a portion of the bovine cGB-8 clone (nucleotides 489-1312 of SEQ ID NO: 9).

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Isolation of cDNA Fragments Encoding Human cGB-PDE

Three human cDNA libraries (two glioblastoma and one lung) in the vector lambda Zap were probed with the bovine cGB-PDE sequence. The PCR-generated clone corresponding to nucleotides 484-1312 of SEQ ID NO: 9 which is described in Example 1 was digested with *Eco*RI and *Sal*I and the resulting 0.8 kb cDNA insert was isolated and purified by agarose gel electrophoresis. The fragment was labelled with radioactive nucleotides using a random primed DNA labelling kit (Boehringer).

The cDNA libraries were plated on 150 mm petri plates at a density of approximately 50,000 plaques per plate. Duplicate nitrocellulose filter replicas were prepared. The prehybridization buffer was 3X SSC, 0.1% sarkosyl, 10X Denhardt's, 20mM sodium phosphate (pH 6.8) and 50 μ g/ml salmon testes DNA. Prehybridization was carried out at 65°C for a minimum of 30 minutes. Hybridization was carried out at 65°C overnight in buffer of the same composition with the addition of 1-5x10° cpm/ml of probe. The filters were washed at 65°C in 2X SSC, 0.1% SDS. Hybridizing plaques were detected by autoradiography. The number of cDNAs that hybridized to the bovine probe and the number of cDNAs screened are indicated in Table 2 below.

20	Table 2					
	cDNA Library	Type	Positive Plaques	Plaques Screened		
	Human SW 1088 glioblastoma	dT-primed	1	1.5x10 ⁶		
	Human lung	dT-primed	2	1.5x10 ⁶		
25	Human SW 1088 glioblastoma	dT-primed	4	1.5x10 ⁶		

Plasmids designated cgbS2.1, cgbS3.1, cgbL23.1, cgbL27.1 and cgbS27.1 were excised in vivo from the lambda Zap clones and sequenced.

Clone cgbS3.1 contains 2060 bp of a PDE open reading frame followed by a putative intron. Analysis of clone cgbS2.1 reveals that it corresponds to clone

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cgbS3.1 positions 664 to 2060 and extends the PDE open reading frame an additional 585 bp before reading into a putative intron. The sequences of the putative 5' untranslated region and the protein encoding portions of the cgbS2.1 and cgbS3.1 clones are set out in SEQ ID NOs: 11 and 12, respectively. Combining the two cDNAs yields a sequence containing approximately 2.7 kb of an open reading encoding a PDE. The three other cDNAs did not extend any further 5' or 3' than cDNA cgbS3.1 or cDNA cgbS2.1.

To isolate additional cDNAs, probes specific for the 5' end of clone cgbS3.1 and the 3' end of clone cgbS2.1 were prepared and used to screen a SW1088 glioblastoma cDNA library and a human aorta cDNA library. A 5' probe was derived from clone cgbS3.1 by PCR using the primers cgbS3.1S311 and cgbL23.1A1286 whose sequences are set out in SEQ ID NOs: 8 and 9, respectively, and below.

Primer cgbS3.1S311 (SEQ ID NO: 13)

5' GCCACCAGAGAAATGGTC 3'

Primer cgbL23.1A1286 (SEQ ID NO: 14)

5' ACAATGGGTCTAAGAGGC 3'

The PCR reaction was carried out in a 50 ul reaction volume containing 50 pg cgbS3.1 cDNA, 0.2mM dNTP, 10 ug/ml each primer, 50 mM KCl, 10mM Tris-HCl pH 8.2, 1.5mM MgCl₂ and Taq polymerase. After an initial four minute denaturation at 94°C, 30 cycles of one minute at 94°C, two minutes at 50°C and four minutes at 72°C were carried out. An approximately 0.2 kb fragment was generated by the PCR reaction which corresponded to nucleotides 300-496 of clone cgbS3.1.

A 3' probe was derived from cDNA cgbS2.1 by PCR using the oligos cgbL23.1S1190 and cgbS2.1A231 whose sequences are set out below.

Primer cgbL23.1S1190 (SEQ ID NO: 15)

5' TCAGTGCATGTTTGCTGC 3'

Primer cgbS2.1A231 (SEQ ID NO: 16)

5' TACAAACATGTTCATCAG 3'

The PCR reaction as carried out similarly to that described above for generating the 5' probe, and yielded a fragment of approximately 0.8kb corresponding to nucleotides

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1358-2139 of cDNA cgbS2.1. The 3' 157 nucleotides of the PCR fragment (not shown in SEQ ID NO: 12) are within the presumptive intron.

The two PCR fragments were purified and isolated by agarose gel electrophoresis, and were labelled with radioactive nucleotides by random priming. A random-primed SW1088 glioblastoma cDNA library (1.5x10⁶ plaques) was screened with the labelled fragments as described above, and 19 hybridizing plaques were isolated. An additional 50 hybridizing plaques were isolated from a human aorta cDNA library (dT and random primed, Clontech, Palo Alto, CA).

Plasmids were excised in vivo from some of the positive lambda Zap clones and sequenced. A clone designated cgbS53.2, the sequence of which is set out in SEQ ID NO: 17, contains an approximately 1.1 kb insert whose sequence overlaps the last 61 bp of cgbS3.1 and extends the open reading frame an additional 135 bp beyond that found in cgbS2.1. The clone contains a termination codon and approximately 0.3 kB of putative 3' untranslated sequence.

Generation of a Composite cDNA Encoding Human cGB-PDE

Clones cgbS3.1, cgbS2.1 and cgbS53.2 were used as described in the following paragraphs to build a composite cDNA that contained a complete human cGB-PDE opening reading frame. The composite cDNA is designated cgbmet156-2 and was inserted in the yeast ADH1 expression vector pBNY6N.

First, a plasmid designated cgb stop-2 was generated that contained the 3' end of the cGB-PDE open reading frame. A portion of the insert of the plasmid was generated by PCR using clone cgbS53.2 as a template. The PCR primers utilized were cgbS2.1S1700 and cgbstop-2.

Primer cgbS2.1S1700 (SEQ ID NO: 18)

5' TITGGAAGATCCTCATCA 3'

Primer cgbstop-2 (SEQ ID NO: 19)

5' ATGTCTCGAGTCAGTTCCGCTTGGCCTG 3'

The PCR reaction was carried out in 50 ul containing 50 pg template DNA, 0.2mM dNTPs, 20mM Tris-HCl pH 8.2, 10mM KC1, 6mM (NH₄)₂SO₄, 1.5mM MgCl₂, 0.1% Triton-X-100, 500ng each primer and 0.5 units of Pfu polymerase (Stratagene). The reaction was heated to 94°C for 4 minutes and then 30 cycles of 1 minute at 94°C, 2 minutes at 50°C and four minutes at 72°C were performed. The polymerase

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was added during the first cycle at 50°C. The resulting PCR product was phenol/chloroform extracted, chloroform extracted, ethanol precipitated and cut with the restriction enzymes *BclI* and *XhoI*. The restriction fragment was purified on an agarose gel and eluted.

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This fragment was ligated to the cDNA cgbS2.1 that had been grown in dam E. coli, cut with the restriction enzymes BclI and XhoI, and gel-purified using the Promega magic PCR kit. The resulting plasmid was sequenced to verify that cgbstop-2 contains the 3' portion of the cGB-PDE open reading frame.

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Second, a plasmid carrying the 5' end of the human cGB-PDE open reading frame was generated. Its insert was generated by PCR using clone cgbS3.1 as a template. PCR was performed as described above using primers cgbmet156 and cgbS2.1A2150.

Primer cgbmet156 (SEQ ID NO: 20)

5' TACAGAATTCTGACCATGGAGCGGGCCGGC 3'

Primer cgbS2.1A2150 (SEQ ID NO: 21)

5' CATTCTAAGCGGATACAG 3'

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The resulting PCR fragment was phenol/choloform extracted, choloform extracted, ethanol precipitated and purified on a Sepharose CL-6B column. The fragment was cut with the restriction enzymes *EcoRV* and *EcoRI*, run on an agarose gel and purified by spinning through glass wool. Following phenol/chloroform extraction, chloroform extraction and ethanol precipitation, the fragment was ligated into *EcoRI/EcoRV* digested BluescriptII SK(+) to generate plasmid cgbmet156. The DNA sequence of the insert and junctions was determined. The insert contains a new *EcoRI* site and an additional 5 nucleotides that together replace the original 155 nucleotides 5' of the initiation codon. The insert extends to an *EcoRV* site beginning 531 nucleotides from the initiation codon.

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The 5' and 3' portions of the cGB-PDE open reading frame were then assembled in vector pBNY6a. The vector pBNY6a was cut with *EcoRI* and *XhoI*, isolated from a gel and combined with the agarose gel purified *EcoRI/EcoRV* fragment from cgbmet156 and the agarose gel purified *EcoRV/XhoI* fragment from cgbstop-2. The junctions of the insert were sequenced and the construct was named hcbgmet156-2 6a.

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The cGB-PDE insert from hobgmet156-2 6a was then moved into the expression vector pBNY6n. Expression of DNA inserted in this vector is directed from the yeast ADH1 promoter and terminator. The vector contains the yeast 2 micron origin of replication, the pUC19 origin of replication and an ampicillan resistance gene. Vector pBNY6n was cut with *Eco*RI and *Xho*I and gel-purified. The EcoRI/XhoI insert from hogbmet156-2 6a was gel purified using Promega magic PCR columns and ligated into the cut pBNY6n. All new junctions in the resulting construct, hogbmet156-2 6n, were sequenced. The DNA and deduced amino acid sequences of the insert of hogbmet156-2 6n which encodes a composite human cGB-PDE is set out in SEQ ID NOs: 22 and 23. The insert extends from the first methionine in clone cgbS3.1 (nucleotide 156) to the stop codon (nucleotide 2781) in the composite cDNA. Because the methionine is the most 5' methionine in clone cgbS3.1 and because there are no stop codons in frame with the methionine and upstream of it, the insert in pBNY6n may represent a truncated form of the open reading frame.

Variant cDNAs

Four human cGB-PDE cDNAs that are different from the hcgbmet156-2 6n composite cDNA have been isolated. One cDNA, cgbL23.1, is missing an internal region of hcgbmet156-2 6n (nucleotides 997-1000 to 1444-1447). The exact end points of the deletion cannot be determined from the cDNA sequence at those positions. Three of the four variant cDNAs have 5' end sequences that diverge from the hcgbmet156-2 6n sequence upstream of nucleotide 151 (cDNAs cgbA7f, cgbA5C, cgbI2). These cDNAs presumably represent alteratively spliced or unspliced mRNAs.

Example 7

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The composite human cGB-PDE cDNA construct, hcgbmet156-2 6n, was transformed into the yeast strain YKS45 (ATCC 74225) (MATα his3 trp1 ura3 leu3 pde1::HIS3 pde2::TRP1) in which two endogenous PDE genes are deleted. Transformants complementing the leu deficiency of the YKS45 strain were selected and assayed for cGB-PDE activity. Extracts from cells bearing the plasmid hcgbmet156-2 6n were determined to display cyclic GMP-specific phosphodiesterase activity by the assay described below.

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One liter of YKS45 cells transformed with the plasmid cgbmet156-2 6n and grown in SC-leu medium to a density of 1-2x10⁷ cells/ml was harvested by centrifugation, washed once with deionized water, frozen in dry ice/ethanol and stored at -70°C. Cell pellets (1-1.5 ml) were thawed on ice in the presence of an equal volume of 25mM Tris-Cl (pH 8.0)/5mM EDTA/5mM EGTA/1mM ophenanthroline/0.5mM AEBSF (Calbiochem)/0.1% β-mercaptoethanol and 10 ug/ml each of aprotinin, leupeptin, and pepstatin A. The thawed cells were added to 2 ml of acid-washed glass beads (425-600 µM, Sigma) in 15 ml Corex tube. Cells were broken with 4 cycles consisting of a 30 second vortexing on setting 1 followed by a 60 second incubation on ice. The cell lysate was centrifuged at 12,000 x g for 10 minutes and the supernatant was passed through a 0.8 μ filter. The supernatant was assayed for cGMP PDE activity as follows. Samples were incubated for 20 minutes at 30°C in the presence of 45mM Tris-Cl (pH 8.0), 2mM EGTA, 1mM EDTA, 0.2mg/ml BSA, 5mM MgCl₂, 0.2mM o-phenanthroline, 2ug/ml each of pepstatin A, leupeptin, and aprotinin, 0.1mM AEBSF, 0.02% β -mercaptoethanol and 0.1mM [3H]cGMP as substrate. [14C]-AMP (0.5 nCi/assay) was added as a recovery standard. The reaction was terminated with stop buffer (0.1M ethanolamine pH 9.0, 0.5M ammonium sulfate, 10mM EDTA, 0.05% SDS final concentration). The product was separated from the cyclic nucleotide substrate by chromatography on BioRad Affi-Gel 601. The sample was applied to a column containing approximately 0.25 ml of Affi-Gel 601 equilibrated in column buffer (0.1M ethanolamine pH 9.0 containing 0.5M ammonium sulfate). The column was washed five times with 0.5 ml of column buffer. The product was eluted with four 0.5 ml aliquots of 0.25 acetic acid and mixed with 5 ml Ecolume (ICN Biochemicals). The radioactive product was measured by scintillation counting.

Example 8

Analysis of expression of cGB-PDE mRNA in human tissues was carried out by RNase protection assay.

A probe corresponding to a portion of the putative cGMP binding domain of cGB-PDE (402 bp corresponding to nucleotides 1450 through 1851 of SEQ ID NO: 13) was generated by PCR. The PCR fragment was inserted into the EcoRI

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site of the plasmid pBSII SK(-) to generate the plasmid RP3. RP3 plasmid DNA was linearized with *Xba*I and antisense probes were generated by a modification of the Stratagene T7 RNA polymerase kit. Twenty-five ng of linearized plasmid was combined with 20 microcuries of alpha ³²P rUTP (800 Ci/mmol, 10 mCi/ml), 1X transcription buffer (40mM TrisCl, pH 8, 8mM MgCl₂, 2mM spermidine, 50mM NaCl), 0.25mM each rATP, rGTP and rCTP, 0.1 units of RNase Block II, 5mM DTT, 8μM rUTP and 5 units of T7 RNA Polymerase in a total volume of 5 μl. The reaction was allowed to proceed 1 hour at room temperature and then the DNA template was removed by digestion with RNase free DNase. The reaction was diluted into 100 μl of 40mM TrisCl, pH 8, 6mM MgCl₂ and 10mM NaCl. Five units of RNase-free DNase were added and the reaction was allowed to continue another 15 minutes at 37°C. The reaction was stopped by a phenol extraction followed by a phenol chloroform extraction. One half volume of 7.5M NH₄OAc was added and the probe was ethanol precipitated.

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The RNase protection assays were carried out using the Ambion RNase Protection kit (Austin, TX) and 10 μ g RNA isolated from human tissues by an acid guanidinium extraction method. Expression of cGB-PDE mRNA was easily detected in RNA extracted from skeletal muscle, uterus, bronchus, skin, right saphenous vein, aorta and SW1088 glioblastoma cells. Barely detectable expression was found in RNA extracted from right atrium, right ventricle, kidney cortex, and kidney medulla. Only complete protection of the RP3 probe was seen. The lack of partial protection argues against the cDNA cgbL23.1 (a variant cDNA described in Example 7) representing a major transcript, at least in these RNA samples.

Example 9

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Polyclonal antisera was raised to E. coli-produced fragments of the human cGB-PDE.

A portion of the human cGB-PDE cDNA (nucleotides 1668-2612 of SEQ ID NO: 22, amino acids 515-819 of SEQ ID NO: 23) was amplified by PCR and inserted into the *E. coli* expression vector pGEX2T (Pharmacia) as a BamHI/EcoRI fragment. The pGEX2T plasmid carries an ampicillin resistance gene, an *E. coli* laq I^q gene and a portion of the Schistosoma japonicum glutathione-S-

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transferase (GST) gene. DNA inserted in the plasmid can be expressed as a fusion protein with GST and can then be cleaved from the GST portion of the protein with thrombin. The resulting plasmid, designated cgbPE3, was transformed into *E. coli* strain LE392 (Stratagene). Transformed cells were grown at 37°C to an OD600 of 0.6. IPTG (isopropylthioalactopyranoside) was added to 0.1mM and the cells were grown at 37°C for an additional 2 hours. The cells were collected by centrifugation and lysed by sonication. Cell debris was removed by centrifugation and the supernatant was fractionated by SDS-PAGE. The gel was stained with cold 0.4M KCl and the GST-cgb fusion protein band was excised and electroeluted. The PDE portion of the protein was separated from the GST portion by digestion with thrombin. The digest was fractionated by SDS-PAGE, the PDE protein was electroeluted and injected subcutaneously into a rabbit. The resultant antisera recognizes both the bovine cGB-PDE fragment that was utilized as antigen and the full length human cGB-PDE protein expressed in yeast (see Example 8).

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Example 10

Polynucleotides encoding various cGB-PDE analogs and cGB-PDE fragments were generated by standard methods.

A. cGB-PDE Analogs

All known cGMP-binding PDEs contain two internally homologous tandem repeats within their putative cGMP-binding domains. In the bovine cGB-PDE of the invention, the repeats span at least residues 228-311 (repeat A) and 410-500 (repeat B) of SEQ ID NO: 10. Site-directed mutagenesis of an aspartic acid that is conserved in repeats A and B of all known cGMP-binding PDEs was used to create analogs of cGB-PDE having either Asp-289 replaced with Ala (D289A) or Asp-478 replaced with Ala (D478A). Recombinant wild type (WT) bovine and mutant bovine cGB-PDEs were expressed in COS-7 cells. cGB-PDE purified from bovine lung (native cGB-PDE) and WT cGB-PDE displayed identical cGMP-binding kinetics with a K_d of approximately 2 μ M and a curvilinear dissociation profile (t_{ν_i} = 1.3 hours at 4°C). This curvilinearity may have been due to the presence of distinct high affinity (slow) and low affinity (fast) sites of cGMP binding. The D289A mutant had significantly decreased affinity for cGMP (K_d > 20 μ M) and a single rate of cGMP-

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association ($t_{44} = 0.5$ hours), that was similar to that calculated for the fast site of WT and native cGB-PDE. This suggested the loss of a slow cGMP-binding site in repeat A of this mutant. Conversely, the D478A mutant showed higher affinity for cGMP (K_d of approximately 0.5 μ M) and a single cGMP-dissociation rate ($t_{44} = 2.8$

hours) that was similar to the calculated rate of the slow site of WT and native cGB-PDE. This suggested the loss of a fast site when repeat B was modified. These results indicate that dimeric cGB-PDE possesses two homologous but kinetically distinct cGMP-binding sites, with the conserved aspartic acid being critical for interaction with cGMP at each site. See, Colbran et al., FASEB J., 8: Abstract 2149

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10 (May 15, 1994).

B. Amino-Terminal Truncated cGB-PDE Polypeptides

A truncated human cGB-PDE polypeptide including amino acids 516-875 of SEQ ID NO: 23 was expressed in yeast. A cDNA insert extending from the NcoI site at nucleotide 1555 of SEQ ID NO: 22 through the XhoI site at the 3' end of SEQ ID NO: 22 was inserted into the ADH2 yeast expression vector YEpC-PADH2d [Price et al., Meth. Enzymol., 185: 308-318 (1990)] that had been digested with NcoI and SalI to generate plasmid YEpC-PADH2d HcGB. The plasmid was transformed into spheroplasts of the yeast strain yBJ2-54 (prcl-407 prb1-1122 pep4-3 leu2 trpl ura3-52 Δpde1::URA3, HIS3 Δpde2::TRP1 cir'). The endogenous PDE genes are deleted in this strain. Cells were grown in SC-leu media with 2% glucose to 10⁷ cells/ml, collected by filtration and grown 24 hours in YEP media containing 3% glycerol. Cells were pelleted by centrifugation and stored frozen. Cells were disrupted with glass beads and the cell homogenate was assayed for phosphodiesterase activity essentially as described in Prpic et al., Anal. Biochem., 208: 155-160 (1993). The truncated human cGB-PDE polypeptide exhibited phosphodiesterase activity.

C. Carboxy-Terminal Truncated cGB-PDE Polypeptides

Two different plasmids encoding carboxy-terminal truncated human cGB-PDE polypeptides were constructed.

Plasmid pBJ6-84Hin contains a cDNA encoding amino acids 1-494 of SEQ ID NO: 23 inserted into the NcoI and SalI sites of vector YEpC-PADH2d. The cDNA insert extends from the NcoI site at nucleotide position 10 of SEQ ID NO: 22

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through the HindIII site at nucleotide position 1494 of SEQ ID NO: 22 followed by a linker and the Sall site of YEpC-PADH2d.

Plasmid pBJ6-84Ban contains a cDNA encoding amino acids 1-549 of SEQ ID NO: 23 inserted into the NcoI and SalI sites of vector YEpC-PADH2d. The cDNA insert extends from the NcoI site at nucleotide position 10 of SEQ ID NO: 22 through the BanI site at nucleotide position 1657 of SEQ ID NO: 22 followed by a linker and the SalI site of YEpC-PADH2d.

The trucated cGB-PDE polypeptides are useful for screening for modulators of cGB-PDE activity.

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Example 11

Monoclonal antibodies reactive with human cGB-PDE were generated. Yeast yBJ2-54 containing the plasmid YEpADH2 HcGB (Example 10B) were fermented in a New Brunswick Scientific 10 liter Microferm. The cGB-PDE cDNA insert in plasmid YEpADH2 HcGB extends from the NcoI site at nucleotide 12 of SEQ ID NO: 22 to the XhoI site at the 3' end of SEQ ID NO: 22. An inoculum of 4 x 10° cells was added to 8 liters of media containing SC-leu, 5% glucose, trace metals, and trace vitamins. Fermentation was maintained at 26°C, agitated at 600 rpm with the standard microbial impeller, and aerated with compressed air at 10 volumes per minute. When glucose decreased to 0.3% at 24 hours post-inoculation the culture was infused with 2 liters of 5X YEP media containing 15% glycerol. At 66 hours post-inoculation the yeast from the ferment was harvested by centrifugation at 4,000 x g for 30 minutes at 4°C. Total yield of biomass from this fermentation approached 350 g wet weight.

Human cGB-PDE enzyme was purified from the yeast cell pellet. Assays for PDE activity using 1 mM cGMP as substrate was employed to follow the chromatography of the enzyme. All chromatographic manipulations were performed at 4°C.

Yeast (29g wet weight) were resuspended in 70ml of buffer A (25mM Tris pH 8.0, 0.25mM DTT, 5mM MgCl₂, 10μ M ZnSO⁴, 1mM benzamidine) and lysed by passing through a microfluidizer at 22-24,000 psi. The lysate was centrifuged at $10,000 \times g$ for 30 minutes and the supernatant was applied to a 2.6 x

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28 cm column containing Pharmacia Fast Flow Q anion exchange resin equilibrated with buffer B containing 20mM BisTris-propane pH 6.8, 0.25mM DTT, 1mM MgCl₂, and 10μ M ZnSO₄. The column was washed with 5 column volumes of buffer B containing 0.125M NaCl and then developed with a linear gradient from 0.125 to 1.0M NaCl. Fractions containing the enzyme were pooled and applied directly to a 5 x 20 cm column of ceramic hydroxyapatite (BioRad) equilibrated in buffer C containing 20mM BisTris-propane pH 6.8, 0.25mM DTT, 0.25MKCl, 1mM MgCl₂, and 10µM ZnSO₄. The column was washed with 5 column volumes of buffer C and eluted with a linear gradient from 0 to 250mM potassium phosphate in buffer C. The pooled enzyme was concentrated 8-fold by ultrafiltration (YM30 membrane, Amicon). The concentrated enzyme was chromatographed on a 2.6 x 90 cm column of Pharmacia Sephacryl S300 (Piscataway, NJ) equilibrated in 25mM BisTris-propane pH 6.8, 0.25mM DTT, 0.25M NaCl, 1mM MgCl₂, and 20µM ZnSO₄. Approximately 4 mg of protein was obtained. The recombinant human cGB-PDE enzyme accounted for approximately 90% of protein obtained as judged by SDS polyacrylamide gel electrophoresis followed by Coomassie blue staining.

The purified protein was used as an antigen to raise monoclonal antibodies. Each of 19 week old Balb/c mice (Charles River Biotechnical Services, Inc., Wilmington, Mass.) was immunized sub-cutaneously with 50 ug purified human cGB-PDE enzyme in a 200 ul emulsion consisting of 50% Freund's complete adjuvant (Sigma Chemical Co.). Subsequent boosts on day 20 and day 43 were administered in incomplete Freund's adjuvant. A pre-fusion boost was done on day 86 using 50 ug enzyme in PBS. The fusion was performed on day 90.

The spleen from mouse #1817 was removed sterilely and placed in 10ml serum free RPMI 1640. A single-cell suspension was formed and filtered through sterile 70-mesh Nitex cell strainer (Becton Dickinson, Parsippany, New Jersey), and washed twice by centrifuging at 200 g for 5 minutes and resuspending the pellet in 20 ml serum free RPMI. Thymocytes taken from 3 naive Balb/c mice were prepared in a similar manner.

NS-1 myeloma cells, kept in log phase in RPMI with 11% Fetalclone (FBS) (Hyclone Laboratories, Inc., Logan, Utah) for three days prior to fusion, were centrifuged at 200 g for 5 minutes, and the pellet was washed twice as described in

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the foregoing paragraph. After washing, each cell suspension was brought to a final volume of 10 ml in serum free RPMI, and 20 μ l was diluted 1:50 in 1 ml serum free RPMI. 20 μ l of each dilution was removed, mixed with 20 μ l 0.4% trypan blue stain in 0.85% saline (Gibco), loaded onto a hemocytometer (Baxter Healthcare Corp., Deerfield, Illinois) and counted.

Two x 10⁸ spleen cells were combined with 4.0 x 10⁷ NS-1 cells, centrifuged and the supernatant was aspirated. The cell pellet was dislodged by tapping the tube and 2 ml of 37°C PEG 1500 (50% in 75 mM Hepes, pH 8.0) (Boehringer Mannheim) was added with stirring over the course of 1 minute, followed by adding 14 ml of serum free RPMI over 7 minutes. An additional 16 ml RPMI was added and the cells were centrifuged at 200 g for 10 minutes. After discarding the supernatant, the pellet was resuspended in 200 ml RPMI containing 15% FBS, 100 μM sodium hypoxanthine, 0.4 μM aminopterin, 16 μM thymidine (HAT) (Gibco), 25 units/ml IL-6 (Boehringer Mannheim) and 1.5 x 10⁶ thymocytes/ml. The suspension was first placed in a T225 flask (Corning, United Kingdom) at 37°C for two hours before being dispensed into ten 96-well flat bottom tissue culture plates (Corning, United Kingdom) at 200 μl/well. Cells in plates were fed on days 3, 4, 5 post fusion day by aspirating approximately 100 μl from each well with an 20 G needle (Becton Dickinson), and adding 100 μl/well plating medium described above except containing 10 units/ml IL-6 and lacking thymocytes.

The fusion was screened initially by ELISA. Immulon 4 plates (Dynatech) were coated at 4°C overnight with purified recombinant human cGB-PDE enzyme (100ng/well in 50mM carbonate buffer pH9.6). The plates were washed 3X with PBS containing 0.05% Tween 20 (PBST). The supernatants from the individual hybridoma wells were added to the enzyme coated wells (50 μ l/well). After incubation at 37°C for 30 minutes, and washing as above, 50 μ l of horseradish peroxidase conjugated goat anti-mouse IgG(fc) (Jackson ImmunoResearch, West Grove, Pennsylvania) diluted 1:3500 in PBST was added. Plates were incubated as above, washed 4X with PBST and 100 μ l substrate consisting of 1 mg/ml o-phenylene diamine (Sigma) and 0.1 μ l/ml 30% H₂O₂ in 100 mM citrate, pH 4.5, was added. The color reaction was stopped in 5 minutes with the addition of 50 μ l of 15% H₂SO₄. A₄₉₀ was read on a plate reader (Dynatech).

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Wells C5G, E4D, F1G, F9H, F11G, J4A, and J5D were picked and renamed 102A, 102B, 102C, 102D, 102E, 102F, and 102G respectively, cloned two or three times, successively, by doubling dilution in RPMI, 15% FBS, 100µM sodium hypoxanathine, 16µM thymidine, and 10 units/ml IL-6. Wells of clone plates were scored visually after 4 days and the number of colonies in the least dense wells were recorded. Selected wells of the each cloning were tested by ELISA.

The monoclonal antibodies produced by above hybridomas were isotyped in an ELISA assay. Results showed that monoclonal antibodies 102A to 102E were IgG1, 102F was IgG2b and 102G was IgG2a.

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All seven monoclonal antibodies reacted with human cGS-PDE as determined by Western analysis.

Example 12

Developing modulators of the biological activities of specific PDEs requires differentiating PDE isozymes present in a particular assay preparation. The classical enzymological approach of isolating PDEs from natural tissue sources and studying each new isozyme is hampered by the limits of purification techniques and the inability to definitively assess whether complete resolution of a isozyme has been achieved. Another approach has been to identify assay conditions which might favor the contribution of one isozyme and minimize the contribution of others in a Still another approach has been the separation of PDEs by preparation. immunological means. Each of the foregoing approaches for differentiating PDE isozymes is time consuming and technically difficult. As a result many attempts to develop selective PDE modulators have been performed with preparations containing more than one isozyme. Moreover, PDE preparations from natural tissue sources are susceptible to limited proteolysis and may contain mixtures of active proteolytic products that have different kinetic, regulatory and physiological properties than the full length PDEs.

Recombinant cGB-PDE polypeptide products of the invention greatly facilitate the development of new and specific cGB-PDE modulators. The use of human recombinant enzymes for screening for modulators has many inherent advantages. The need for purification of an isozyme can be avoided by expressing

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it recombinantly in a host cell that lacks endogenous phosphodiesterase activity (e.g., yeast strain YKS45 deposited as ATCC 74225). Screening compounds against human protein avoids complications that often arise from screening against non-human protein where a compound optimized on a non-human protein may fail to be specific for or react with the human protein. For example, a single amino acid difference between the human and rodent 5HT_{1B} serotonin receptors accounts for the difference in binding of a compound to the receptors. [See Oskenberg et al., Nature, 360: 161-163 (1992)]. Once a compound that modulates the activity of the cGB-PDE is discovered, its selectivity can be evaluated by comparing its activity on the cGB-PDE to its activity on other PDE isozymes. Thus, the combination of the recombinant cGB-PDE products of the invention with other recombinant PDE products in a series of independent assays provides a system for developing selective modulators of cGB-PDE. Selective modulators may include, for example, antibodies and other proteins or peptides which specifically bind to the cGB-PDE or cGB-PDE nucleic acid, oligonucleotides which specifically bind to the cGB-PDE (see Patent Cooperation Treaty International Publication No. WO93/05182 published March 18, 1993 which describes methods for selecting oligonucleotides which selectively bind to target biomolecules) or cGB-PDE nucleic acid (e.g., antisense oligonucleotides) and other non-peptide natural or synthetic compounds which specifically bind to the cGB-PDE or cGB-PDE nucleic acid. Mutant forms of the cGB-PDE which alter the enzymatic activity of the cGB-PDE or its localization in a cell are also contemplated. Crystallization of recombinant cGB-PDE alone and bound to a modulator, analysis of atomic structure by X-ray crystallography, and computer modelling of those structures are methods useful for designing and optimizing non-peptide selective modulators. See, for example, Erickson et al., Ann. Rep. Med. Chem., 27: 271-289 (1992) for a general review of structure-based drug design.

Targets for the development of selective modulators include, for example: (1) the regions of the cGB-PDE which contact other proteins and/or localize the cGB-PDE within a cell, (2) the regions of the cGB-PDE which bind substrate, (3) the allosteric cGMP-binding site(s) of cGB-PDE, (4) the metal-binding regions of the cGB-PDE, (5) the phosphorylation site(s) of cGB-PDE and (6) the regions of the cGB-PDE which are involved in dimerization of cGB-PDE subunits.

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While the present invention has been described in terms of specific embodiments, it is understood that variations and modifications will occur to those skilled in the art. Accordingly, only such limitations as appear in the appended claims should be placed on the invention.

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SEQUENCE LISTING

- (1) GENERAL INFORMATION:
 - (i) APPLICANT: The Board of Regents of the University of Washington
 - (ii) TITLE OF INVENTION: Cyclic GMP-Binding, Cyclic GMP-Specific Phosphodiesterase Materials and Methods
 - (iii) NUMBER OF SEQUENCES: 23
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 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
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 - (vii) PRIOR APPLICATION DATA:
 - (A) APPLICATION NUMBER: US 08/068,051
 - (B) FILING DATE: 27-MAY-1993
 - (viii) ATTORNEY/AGENT INFORMATION:
 - (A) NAME: Noland, Greta E.
 - (B) REGISTRATION NUMBER: 35,302
 - (C) REFERENCE/DOCKET NUMBER: 32083
 - (ix) TELECOMMUNICATION INFORMATION:
 - (A) TELEPHONE: (312) 474-6300
 - (B) TELEFAX: (312) 474-0448
 - (C) TELEX: 25-3856
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

Arg Glu Xaa Asp Ala Asn Arg Ile Asn Tyr Met Tyr Ala Gln Tyr Val 1 5 10 15

Lys Asn Thr Met 20

- (2) INFORMATION FOR SEQ ID NO:2:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 9 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Gln Ser Leu Ala Ala Ala Val Val Pro

- (2) INFORMATION FOR SEQ ID NO:3:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 8 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: peptide
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Phe Asp Asn Asp Glu Gly Glu Gln

- (2) INFORMATION FOR SEQ ID NO:4:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

TTYGAYAAYG AYGARGGNGA RCA

- (2) INFORMATION FOR SEQ ID NO:5:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 base pairs (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA
 - (iv) ANTI-SENSE: YES

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(ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION: 99..2723

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:9

			-		~ m									~ > ~ =		_	.
GGG	AGGG	TCT	CGAG	GCGA	GT T	CTGC	TCCT	C GG	AGGG	AGGG	ACC	CCAG	CTG	GAGT	GGAAA		60
CCA	.GCAC	CAG	CTGA	CCGC	AG A	gaca	.CGCC	G CG	CTGA		TG G et G l					1	13
					Ala										CGA Arg	10	61
	_		_	Arg			_		Tyr						AGG Arg	20	09
		ACC Thr 40	Arg							_						25	57
															TGC Cys	30	05
	Сув	CCC Pro		_											GGA Gly 85	35	53
		ACC Thr														40	01
		GTT Val				-								_	_	44	49
		AAG Lys 120		_	_				_							49	97
		GJY GGG														54	45
		CAC His														59	93
		GGA Gly										_		_		64	11
		AGC Ser														68	39
		GGT Gly 200	-													73	37

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		P As	_				Gly					Phe			CCC Pro	785
	ı Ası		C AAA e Lys			Tyr					Phe					833
			r ACA e Thr		Tyr					Ile					Ile	881
			AGG Arg 265	Glu					Val							929
			AAT Asn	_												977
		Tyr	TTG Leu								_		_			1025
	Glu	_	TCA Ser	_							_	_				1073
	_		TTA Leu					_								1121
			GCT Ala 345													1169
			ATA Ile													1217
			GAG Glu		_									-		1265
			GAT Asp			_				_		_	_			1313
			ATG Met													1361
			TGG Trp 425													1409
	Arg		TTG Leu			Thr					_					1457
Val		_	GTT Val		Gln					Met		_				1505

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	Val					CGC Arg					Phe				1553
					Leu	GGG		_				_		GCA Ala	1601
_	_			Met		AAG Lys	_		Val						1649
			Ser			GAG Glu									1697
		Ala				TCT Ser 540		_					 		1745
	Phe					CTG Leu				_					1793
						CTC									1841
						TGG Trp								-	1889
						AAT Asn									1937
						AAA Lys 620		_						ACG Thr	1985
						CTG Leu									2033
			_			AAC Asn									2081
						CAT His									2129
						AAT Asn									2177
						AAG Lys 700									2225
				Asp		GCA Ala									2273
-			Ile			AAT Asn						_			2321

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					_			ATG Met 750							_	2369	€
								CAA Gln								2417	7
								AGA Arg								246	5
								GAG Glu					_			2513	3
								ATC Ile								2561	L
								TTC Phe 830								2609	•
	Asn							CTT Leu							_	2657	7
Leu					Ser			ACC Thr								2709	5
GTT Val 870	_			Thr		TAGC	CAGG	TG T	atca	GATG	a Gi	gagi	GTG1	•		2753	}
GCTC	AGCT	CA G	TCCT	CTGC	A AC	ACCA	TGAA	GCT	AGGC	TTA	CCAG	CTTA	I TA	CCTC	CAGTT	2813	3
GACT	TTAA	AA' A	ACTG	GCAT	A AA	GCAC	TAGT	CAG	CATC	TAG	TTCT	AGCT	TG A	CCAG	TGAAG	2873	3
AGTA	GAAC	AC C	ACCA	CAGT	C AG	GGTG	CAGA	GCA	GTTG	GCA	GTCT	CCTT	TC C	AACC	CAGAC	2933	ţ
TGGT	GAAT	TT A	AAGA	AGAG	CAG	TCGT	CGTT	TAT	atct	CTG	TCTT	TTCC	TA A	cced	GGTGT	2993	ļ
GGAA'	TCTC	TA A	G A GG:	AGAG	A GA	GATC	TGGA	CCA	CAGG	TCC	AATG	CGCT	CT G	TCCI	CTCAG	3053	ļ
CTGC:	TTCC	CC C	actg:	TGCT	G TG	ACCT	CTCA	ATC	TGAG	AAA	CGTG	TAAG	GA A	GGTI	TCAGC	3113	į
GAAT:	rccc'	TT T	AAAA!	rgtg	CA	GACA	GTAG	CTT	CTTG	GGC	CGGG	TTGT	TC C	CGCA	GCTCC	3173	ļ
CCAT	CTGT	rt G	rtgt(CTATO	TT	GGCT	GAAA	GAG	GCTT	TGC	TGTA	CCTG	CC A	CACT	CTCCT	3233	ļ
GGAT	CCT	T C	CAGTI	AGCTO	AT	CAAA	AAAA	AGG	ATGT	GAA .	ATTC	TCGT	GT G	ACTT	TTTAG	3293	}
) A A A A	GAAI	AG TO	BACC	CCGAC	GA!	rcgg:	rgtg	GAT	TCAC	TAG	TTGT	CCAC	AG A	TGAT	CTGTT	3353	J
TAGTI	rtct?	AG AZ	ATTT	CCAP	GA:	rgatz	ACAC	TCC	rccc	TAG	TCTA	GGGG	TC A	GACC	CTGTA	3413	ļ
TGGT	GCT	T G	ACCCI	TGAG	GA	ACTT(CTCT	CTT	rgca:	TGA	CATT:	AGCC	AT A	GAAC	TGTTC	3473	j
TTGT	CAAJ	AT AC	CACAC	CTC	TAT	rgcac	-CTT	GCA	GAA	ACA	CTTT.	AAAA	AC A	CAAC	TATCA	3533	ļ
CCTAI	GTT?	T TC	TGAT	TACA	GA)	AGTT!	ATCC	CTA	CTCA	CTG	TAAA	CATA	AA C	AAAG	cccc	3593	ļ
CAAAC	TTC	IA AI	AGTI	rgtg1	GTO	GTG!	AGAA	ACTO	CAA	GTT '	TTCA'	TCTC	CA G	agat	AGCTA	3653	,
raggi	AATI	A GT	GGGA	TGTT	TCI	[GAAZ	CTT	TTA	יבבב	מבי	тстт	TTAC	AT A	TATG	TTAAC	3713	1

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TGTTTTCTTA	TGAGCACTAT	GGTTTGTTTT	TTTTTTTTT	TGCTCTGCTT	TGACTTGCCC	3773
TTTTCACTCA	ATTATCTTGG	CAGTTTTTCT	AAATGACTTG	CACAGACTTC	TCCTGTACTT	3833
CATGGCTGTG	CAGTGTTCCA	TGCTGTGAAG	GCACCATCGT	GTATTAAATC	AGTTCCCTGG	3893
TCACACATAG	GTGAGCTGGT	TGGAAATTTT	TACCATTAAA	AAACCACTTT	CCCACATTGA	3953
TGCTTTCTAA	TCTGGCACAG	GATGCTTCTT	TTTTTCCCCT	TTTTCTCTGT	TTAATTATTG	4013
GAAATGGGAT	CTGTGGGATC	CTCGTTCCCT	GGCACCTAGC	TGCTCTCAAC	GTGGCCTGTG	4073
GCCAGCAGCA	TTGGCTAGAC	CTGGGGGCTT	GTTGGGAACG	GAGACCCTCT	GCCCTGCCCC	4133
TGGCCTGCTG	ACAAGGACCT	GCATTTTGCT	GAGCTCCCAG	TGACCCTGGT	GTTTAATTGT	4193
TAACCATTGA	AAAAAATCAA	ACTATAGTTT	ATTTACAATG	TTGTGTTAAT	TTCGGGTGTA	4253
CAGCAAAGTG	ACTCAGTGGT	CAAGTACATT	TAAAACACTG	GGCATACTCT	CTCCCTCTCC	4313
TTGTGTACCT	GGTTGGTATT	TCCAGAAACC	ATGCTCTTGT	CTGTCCTGTA	GTTTTGGAAG	4373
CGCTTTCTCT	TTGAAGACTG	CCTTCTCTCC	TGTGTCTGCC	CTACATGGAC	TAGTTCGTTT	4433
ATTGTCCTAC	ATGGCTTTGC	TTCCATGTTC	CTCTCAACTT	T		4474

(2) INFORMATION FOR SEQ ID NO:10:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 875 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

Met Glu Arg Ala Gly Pro Gly Cys Arg Ala Ala Ala Thr Ala Met Gly 1 10 15 Pro Gly Leu Gly Arg Ser Val Ala Gly Arg Ser Leu Gly Leu Tyr Leu Leu Tyr Phe Val Arg Lys Gly Thr Arg Glu Met Val Asn Ala Trp Phe Ala Glu Arg Val His Thr Ile Pro Val Cys Lys Glu Gly Ile Lys Gly His Thr Glu Ser Cys Ser Cys Pro Leu Gln Pro Ser Pro Arg Ala Glu

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Ser Ser Val Pro Gly Thr Pro Thr Arg Lys Ile Ser Ala Ser Glu Phe

Asp Arg Pro Leu Arg Pro Ile Val Ile Lys Asp Ser Glu Gly Thr Val 105 100

Ser Phe Leu Ser Asp Ser Asp Lys Lys Glu Gln Met Pro Leu Thr Ser 115

Pro Arg Phe Asp Asn Asp Glu Gly Asp Gln Cys Ser Arg Leu Leu Glu 140 130

Leu Val Lys Asp Ile Ser Ser His Leu Asp Val Thr Ala Leu Cys His 150 155 145

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Lys Ile Phe Leu His Ile His Gly Leu Ile Ser Ala Asp Arg Tyr Ser 175 165 Leu Phe Leu Val Cys Glu Asp Ser Ser Asn Asp Lys Phe Leu Ile Ser 180 Arg Leu Phe Asp Val Ala Glu Gly Ser Thr Leu Glu Glu Ala Ser Asn 195 Asn Cys Ile Arg Leu Glu Trp Asn Lys Gly Ile Val Gly His Val Ala 210 Ala Phe Gly Glu Pro Leu Asn Ile Lys Asp Ala Tyr Glu Asp Pro Arg 225 230 Phe Asn Ala Glu Val Asp Gln Ile Thr Gly Tyr Lys Thr Gln Ser Ile 245 250 Leu Cys Met Pro Ile Lys Asn His Arg Glu Glu Val Val Gly Val Ala 260 265 Gln Ala Ile Asn Lys Lys Ser Gly Asn Gly Gly Thr Phe Thr Glu Lys Asp Glu Lys Asp Phe Ala Ala Tyr Leu Ala Phe Cys Gly Ile Val Leu His Asn Ala Gln Leu Tyr Glu Thr Ser Leu Leu Glu Asn Lys Arg Asn 305 Gln Val Leu Leu Asp Leu Ala Ser Leu Ile Phe Glu Glu Gln Gln Ser Leu Glu Val Ile Leu Arg Lys Ile Ala Ala Thr Ile Ile Ser Pro Met Gln Val Gln Lys Cys Thr Ile Phe Ile Val Asp Glu Asp Cys Ser Asp 360 Ser Phe Ser Ser Val Phe His Met Glu Cys Glu Glu Leu Glu Lys Ser 370 375 Ser Asp Thr Leu Thr Arg Glu Arg Asp Ala Thr Arg Ile Asn Tyr Met 385 Tyr Ala Gln Tyr Val Lys Asn Thr Met Glu Pro Leu Asn Ile Pro Asp Val Ser Lys Asp Lys Arg Phe Pro Trp Thr Asn Glu Asn Met Gly Asn Ile Asn Gln Gln Cys Ile Arg Ser Leu Leu Cys Thr Pro Ile Lys Asn Gly Lys Lys Asn Lys Val Ile Gly Val Cys Gln Leu Val Asn Lys Met Glu Glu Thr Thr Gly Lys Val Lys Ala Phe Asn Arg Asn Asp Glu Gln 465 470 Phe Leu Glu Ala Phe Val Ile Phe Cys Gly Leu Gly Ile Gln Asn Thr 485 Gln Met Tyr Glu Ala Val Glu Arg Ala Met Ala Lys Gln Met Val Thr 500

850

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Leu Glu Val Leu Ser Tyr His Ala Ser Ala Ala Glu Glu Glu Thr Arg 515 Glu Leu Gln Ser Leu Ala Ala Ala Val Val Pro Ser Ala Gln Thr Leu 530 **535** 540 Lys Ile Thr Asp Phe Ser Phe Ser Asp Phe Glu Leu Ser Asp Leu Glu 545 550 Thr Ala Leu Cys Thr Ile Arg Met Phe Thr Asp Leu Asn Leu Val Gln 565 570 575 Asn Phe Gln Met Lys His Glu Val Leu Cys Lys Trp Ile Leu Ser Val Lys Lys Asn Tyr Arg Lys Asn Val Ala Tyr His Asn Trp Arg His Ala 600 Phe Asn Thr Ala Gln Cys Met Phe Ala Ala Leu Lys Ala Gly Lys Ile Gln Lys Arg Leu Thr Asp Leu Glu Ile Leu Ala Leu Leu Ile Ala Ala 630 Leu Ser His Asp Leu Asp His Arg Gly Val Asn Asn Ser Tyr Ile Gln 645 650 Arg Ser Glu His Pro Leu Ala Gln Leu Tyr Cys His Ser Ile Met Glu His His His Phe Asp Gln Cys Leu Met Ile Leu Asn Ser Pro Gly Asn 675 680 Gln Ile Leu Ser Gly Leu Ser Ile Glu Glu Tyr Lys Thr Thr Leu Lys 690 695 700 Ile Ile Lys Gln Ala Ile Leu Ala Thr Asp Leu Ala Leu Tyr Ile Lys 705 710 720 715 Arg Arg Gly Glu Phe Phe Glu Leu Ile Met Lys Asn Gln Phe Asn Leu 725 730 735 Glu Asp Pro His Gln Lys Glu Leu Phe Leu Ala Met Leu Met Thr Ala Cys Asp Leu Ser Ala Ile Thr Lys Pro Trp Pro Ile Gln Gln Arg Ile Ala Glu Leu Val Ala Thr Glu Phe Phe Asp Gln Gly Asp Arg Glu Arg 770 Lys Glu Leu Asn Ile Glu Pro Ala Asp Leu Met Asn Arg Glu Lys Lys 785 790 795 Asn Lys Ile Pro Ser Met Gln Val Gly Phe Ile Asp Ala Ile Cys Leu 805 815 Gln Leu Tyr Glu Ala Leu Thr His Val Ser Glu Asp Cys Phe Pro Leu 820 Leu Asp Gly Cys Arg Lys Asn Arg Gln Lys Trp Gln Ala Leu Ala Glu 835 840 Gln Gln Glu Lys Thr Leu Ile Asn Gly Glu Ser Ser Gln Thr Asn Arg

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Gln Gln Arg Asn Ser Val Ala Val Gly Thr Val 870 865

(2) INFORMATION FOR SEQ ID NO:11:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 2060 base pairs (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

GCGGCGCGCG	C TCCGGCCGCT	TTGTCGAAAG	CCGGCCCGAC	TGGAGCAGGA	CGAAGGGGGA	60
GGGTCTCGAC	GCCGAGTCCI	GTTCTTCTGA	GGGACGGACC	CCAGCTGGGG	TGGAAAAGCA	120
GTACCAGAGA	GCCTCCGAGG	CGCGCGGTGC	CAACCATGGA	GCGGGCCGGC	CCCAGCTTCG	180
GGCAGCAGCG	ACAGCAGCAG	CAGCCCCAGC	AGCAGAAGCA	GCAGCAGAGG	GATCAGGACT	240
CGGTCGAAGC	ATGGCTGGAC	GATCACTGGG	ACTITACCTT	CTCATACTTT	GTTAGAAAAG	300
CCACCAGAGA	AATGGTCAAT	GCATGGTTTG	CTGAGAGAGT	TCACACCATC	CCTGTGTGCA	360
AGGAAGGTAT	CAGAGGCCAC	ACCGAATCTT	GCTCTTGTCC	CTTGCAGCAG	AGTCCTCGTG	420
CAGATAACAG	TGTCCCTGGA	ACACCAACCA	GGAAAATCTC	TGCCTCTGAA	TTTGACCGGC	480
CTCTTAGACC	CATTGTTGTC	AAGGATTCTG	AGGGAACTGT	GAGCTTCCTC	TCTGACTCAG	540
AAAAGAAGGA	ACAGATGCCT	CTAACCCCTC	CAAGGTTTGA	TCATGATGAA	GGGGACCAGT	600
GCTCAAGACT	CTTGGAATTA	GTGAAGGATA	TTTCTAGTCA	TTTGGATGTC	ACAGCCTTAT	660
GTCACAAAAT	TTTCTTGCAT	ATCCATGGAC	TGATATCTGC	TGACCGCTAT	TCCCTGTTCC	720
TTGTCTGTGA	AGACAGCTCC	AATGACAAGT	TTCTTATCAG	CCGCCTCTTT	GATGTTGCTG	780
AAGGTTCAAC	ACTGGAAGAA	GTTTCAAATA	ACTGTATCCG	CTTAGAATGG	AACAAAGGCA	840
TTGTGGGACA	TGTGGCAGCG	CTTGGTGAGC	CCTTGAACAT	CAAAGATGCA	TATGAGGATC	900
CTCGGTTCAA	TGCAGAAGTT	GACCAAATTA	CAGGCTACAA	GACACAAAGC	ATTCTTTGTA	960
TGCCAATTAA	GAATCATAGG	GAAGAGGTTG	TTGGTGTAGC	CCAGGCCATC	AACAAGAAAT	1020
CAGGAAACGG	TGGGACATTT	ACTGAAAAAG	ATGAAAAGGA	CTTTGCTGCT	TATTTGGCAT	1080
TTTGTGGTAT	TGTTCTTCAT	AATGCTCAGC	TCTATGAGAC	TTCACTGCTG	GAGAACAAGA	1140
GAAATCAGGT	GCTGCTTGAC	CTTGCTAGTT	TAATTTTTGA	AGAACAACAA	TCATTAGAAG	1200
TAATTTTGAA	GAAAATAGCT	GCCACTATTA	TCTCTTTCAT	GCAAGTGCAG	AAATGCACCA	1260
TTTTCATAGT	GGATGAAGAT	TGCTCCGATT	CTTTTTCTAG	TGTGTTTCAC	ATGGAGTGTG	1320
aggaattaga	AAAATCATCT	GATACATTAA	CAAGGGAACA	TGATGCAAAC	AAAATCAATT	1380
ACATGTATGC	TCAGTATGTC	aaaaatacta	TGGAACCACT	TTATATCCCA	GATGTCAGTA	1440
AGGATAAAAG	ATTTCCCTGG	ACAACTGAAA	ATACAGGAAA	TGTAAACCAG	CAGTGCATTA	1500

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GAAGTTTGCT	TTGTACACCT	ATAAAAAATG	GAAAGAAGAA	TAAAGTTATA	GGGGTTTGCC	1560
AACTTGTTAA	TAAGATGGAG	GAGAATACTG	GCAAGGTTAA	GCCTTTCAAC	CGAAATGACG	1620
AACAGTTTCT	GGAAGCTTTT	GTCATCTTTT	GTGGCTTGGG	GATCCAGAAC	ACGCAGATGT	1680
ATGAAGCAGT	GGAGAGAGCC	ATGGCCAAGC	AAATGGTCAC	ATTGGAGGTT	CTGTCGTATC	1740
ATGCTTCAGC	AGCAGAGGAA	GAAACAAGAG	AGCTACAGTC	GTTAGCGGCT	GCTGTGGTGC	1800
CATCTGCCCA	GACCCTTAAA	ATTACTGACT	TTAGCTTCAG	TGACTTTGAG	CTGTCTGATC	1860
TGGAAACAGC	ACTGTGTACA	ATTCGGATGT	TTACTGACCT	CAACCTTGTG	CAGAACTTCC	1920
AGATGAAACA	TGAGGTTCTT	TGCAGATGGA	TTTTAAGTGT	TAAGAAGAAT	TATCGGAAGA	1980
ATGTTGCCTA	TCATAATTGG	AGACATGCCT	TTAATACAGC	TCAGTGCATG	TTTGCTGCTC	2040
TAAAAGCAGG	CAAAATTCAG					2060

(2) INFORMATION FOR SEQ ID NO:12:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1982 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

ACAAAATTTT	CTTGCATATC	CATGGACTGA	TATCTGCTGA	CCGCTATTCC	CTGTTCCTTG	60	
TCTGTGAAGA	CAGCTCCAAT	GACAAGTTTC	TTATCAGCCG	CCTCTTTGAT	GTTGCTGAAG	120	
GTTCAACACT	GGAAGAAGTT	TCAAATAACT	GTATCCGCTT	AGAATGGAAC	AAAGGCATTG	180	
TGGGACATGT	GGCAGCGCTT	GGTGAGCCCT	TGAACATCAA	AGATGCATAT	GAGGATCCTC	240	
GGTTCAATGC	AGAAGTTGAC	CAAATTACAG	GCTACAAGAC	ACAAAGCATT	CTTTGTATGC	300	
CAATTAAGAA	TCATAGGGAA	GAGGTTGTTG	GTGTAGCCCA	GGCCATCAAC	AAGAAATCAG	360	
GAAACGGTGG	GACATTTACT	GAAAAAGATG	AAAAGGACTT	TGCTGCTTAT	TTGGCATTTT	420	
GTGGTATTGT	TCTTCATAAT	GCTCAGCTCT	ATGAGACTTC	ACTGCTGGAG	AACAAGAGAA	480	
ATCAGGTGCT	GCTTGACCTT	GCTAGTTTAA	TTTTTGAAGA	ACAACAATCA	TTAGAAGTAA	540	
TTTTGAAGAA	AATAGCTGCC	ACTATTATCT	CTTTCATGCA	AGTGCAGAAA	TGCACCATTT	600	
TCATAGTGGA	TGAAGATTGC	TCCGATTCTT	TTTCTAGTGT	GTTTCACATG	GAGTGTGAGG	660	
AATTAGAAAA	ATCATCTGAT	ACATTAACAA	GGGAACATGA	TGCAAACAAA	ATCAATTACA	720	
TGTATGCTCA	GTATGTCAAA	AATACTATGG	AACCACTTAA	TATCCCAGAT	GTCAGTAAGG	780	
ATAAAAGATT	TCCCTGGACA	ACTGAAAATA	CAGGAAATGT	AAACCAGCAG	TGCATTAGAA	840	
GTTTGCTTTG	TACACCTATA	AAAAATGGAA	AGAAGAATAA	AGTTATAGGG	GTTTGCCAAC	900	
TTGTTAATAA	GATGGAGGAG	AATACTGGCA	AGGTTAAGCC	TTTCAACCGA	AATGACGAAC	960	
AGTTTCTGGA	AGCTTTTGTC	ATCTTTTGTG	GCTTGGGGAT	CCAGAACACG	CAGATGTATG	1020	

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AAGCAGTGGA	GAGAGCCATG	GCCAAGCAAA	TGGTCACATT	GGAGGTTCTG	TCGTATCATG	1080
CTTCAGCAGC	AGAGGAAGAA	ACAAGAGAGC	TACAGTCGTT	AGCGGCTGCT	GTGGTGCCAT	1140
CTGCCCAGAC	CCTTAAAATT	ACTGACTTTA	GCTTCAGTGA	CTTTGAGCTG	TCTGATCTGG	1200
AAACAGCACT	GTGTACAATT	CGGATGTTTA	CTGACCTCAA	CCTTGTGCAG	AACTTCCAGA	1260
TGAAACATGA	GGTTCTTTGC	AGATGGATTT	TAAGTGTTAA	GAAGAATTAT	CGGAAGAATG	1320
TTGCCTATCA	TAATTGGAGA	CATGCCTTTA	ATACAGCTCA	GTGCATGTTT	GCTGCTCTAA	1380
AAGCAGGCAA	AATTCAGAAC	AAGCTGACTG	ACCTGGAGAT	ACTTGCATTG	CTGATTGCTG	1440
CACTAAGCCA	CGATTTGGAT	CACCGTGGTG	TGAATAACTC	TTACATACAG	CGAAGTGAAC	1500
ATCCACTTGC	CCAGCTTTAC	TGCCATTCAA	TCATGGAACA	CCATCATTTT	GACCAGTGCC	1560
TGATGATTCT	TAATAGTCCA	GGCAATCAGA	TTCTCAGTGG	CCTCTCCATT	GAAGAATATA	1620
AGACCACGTT	GAAAATAATC	AAGCAAGCTA	TTTTAGCTAC	AGACCTAGCA	CTGTACATTA	1680
AGAGGCGAGG	AGAATTTTTT	GAACTTATAA	GAAAAAATCA	ATTCAATTTG	GAAGATCCTC	1740
ATCAAAAGGA	GTTGTTTTTG	GCAATGCTGA	TGACAGCTTG	TGATCTTTCT	GCAATTACAA	1800
AACCCTGGCC	TATTCAACAA	CGGATAGCAG	AACTTGTAGC	AACTGAATTT	TTTGATCAAG	1860
GAGACAGAGA	GAGAAAAGAA	CTCAACATAG	AACCCACTGA	TCTAATGAAC	AGGGAGAAGA	1920
AAAACAAAAT	CCCAAGTATG	CAAGTTGGGT	TCATAGATGC	CATCTGCTTG	CAACTGTATG	1980
AG						1982

(2) INFORMATION FOR SEQ ID NO:13:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 18 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

GCCACCAGAG AAATGGTC

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- (2) INFORMATION FOR SEQ ID NO:14:
 - (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

ACAATGGGTC TAAGAGGC

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(2)	INFORMATION	FOR	SEO	ID	NO:15:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 18 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

TCAGTGCATG TTTGCTGC

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- (2) INFORMATION FOR SEQ ID NO:16:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 18 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:

TACAAACATG TTCATCAG

- (2) INFORMATION FOR SEQ ID NO:17:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1107 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:

•						
GAGACATGCC	TTTAATACAG	CTCAGTGCAT	GTTTGCTGCT	CTAAAAGCAG	GCAAAATTCA	60
GAACAAGCTG	ACTGACCTGG	AGATACTTGC	ATTGCTGATT	GCTGCACTAA	GCCACGATTT	120
GGATCACCGT	GGTGTGAATA	ACTCTTACAT	ACAGCGAAGT	GAACATCCAC	TTGCCCAGCT	180
TTACTGCCAT	TCAATCATGG	AACACCATCA	TTTTGACCAG	TGCCTGATGA	TTCTTAATAG	240
TCCAGGCAAT	CAGATTCTCA	GTGGCCTCTC	CATTGAAGAA	TATAAGACCA	CGTTGAAAAT	300
AATCAAGCAA	GCTATTTTAG	CTACAGACCT	AGCACTGTAC	ATTAAGAGGC	GAGGAGAATT	360
TTTTGAACTT	ATAAGAAAAA	ATCAATTCAA	TTTGGAAGAT	CCTCATCAAA	AGGAGTTGTT	420
TTTGGCAATG	CTGATGACAG	CTTGTGATCT	TTCTGCAATT	ACAAAACCCT	GGCCTATTCA	480
ACAACGGATA	GCAGAACTTG	TAGCAACTGA	ATTTTTTGAT	CAAGGAGACA	GAGAGAGAAA	540
AGAACTCAAC	ATAGAACCCA	CTGATCTAAT	GAACAGGGAG	AAGAAAAACA	AAATCCCAAG	600
TATGCAAGTT	GGGTTCATAG	ATGCCATCTG	CTTGCAACTG	TATGAGGCCC	TGACCCACGT	660

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GTCAGAGGAC	TGTTTCCCTT	TGCTAGATGG	CTGCAGAAAG	AACAGGCAGA	AATGGCAGGC	720
CCTTGCAGAA	CAGCAGGAGA	AGATGCTGAT	TAATGGGGAA	AGCGGCCAGG	CCAAGCGGAA	780
CTGAGTGGCC	TATTTCATGC	AGAGTTGAAG	TTTACAGAGA	TGGTGTGTTC	TGCAATATGC	840
CTAGTTTCTT	ACACACTGTC	TGTATAGTGT	CTGTATTTGG	TATATACTTT	GCCACTGCTG	900
TATTTTTATT	TTTGCACAAC	TTTTGAGAGT	ATAGCATGAA	TGTTTTTAGA	GGACTATTAC	960
ATATTTTTTG	TATATTTGTT	TTATGCTACT	GAACTGAAAG	GATCAACAAC	ATCCACTGTT	1020
AGCACATTGA	TAAAAGCATT	GTTTGTGATA	TTTCGTGTAC	TGCAAAGTGT	ATGCAGTATT	1080
CTTGCACTGA	GCTTTTTTTG	CTTGGGG				1107

(2) INFORMATION FOR SEQ ID NO:18:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 18 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

TTTGGAAGAT CCTCATCA

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- (2) INFORMATION FOR SEQ ID NO:19:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 28 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

ATGTCTCGAG TCAGTTCCGC TTGGCCTG

28

- (2) INFORMATION FOR SEQ ID NO:20:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 30 base pairs (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

TACAGAATTC TGACCATGGA GCGGGCCGGC

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(2) INFORMATION FOR SEQ ID NO:21:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:	
CATTCTAAGC GGATACAG	18
(2) INFORMATION FOR SEQ ID NO:22:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 2645 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: cDNA	
(1x) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 122636	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:22: GAATTCTGAC C ATG GAG CGG GCC GGC CCC AGC TTC GGG CAG CAG CAG Met Glu Arg Ala Gly Pro Ser Phe Gly Gln Gln Arg Gln 1	50
CAG CAG CCC CAG CAG CAG AAG CAG CAG CAG	98
GTC GAA GCA TGG CTG GAC GAT CAC TGG GAC TTT ACC TTC TCA TAC TTT Val Glu Ala Trp Leu Asp Asp His Trp Asp Phe Thr Phe Ser Tyr Phe 35	146
GTT AGA AAA GCC ACC AGA GAA ATG GTC AAT GCA TGG TTT GCT GAG AGA Val Arg Lys Ala Thr Arg Glu Met Val Asn Ala Trp Phe Ala Glu Arg 50 55 60	194
GTT CAC ACC ATC CCT GTG TGC AAG GAA GGT ATC AGA GGC CAC ACC GAA Val His Thr Ile Pro Val Cys Lys Glu Gly Ile Arg Gly His Thr Glu 65 70 75	242
TCT TGC TCT TGT CCC TTG CAG CAG AGT CCT CGT GCA GAT AAC AGT GTC Ser Cys Ser Cys Pro Leu Gln Gln Ser Pro Arg Ala Asp Asn Ser Val 80 85 90	290
CCT GGA ACA CCA ACC AGG AAA ATC TCT GCC TCT GAA TTT GAC CGG CCT Pro Gly Thr Pro Thr Arg Lys Ile Ser Ala Ser Glu Phe Asp Arg Pro 95	338

CTT AGA CCC ATT GTT GTC AAG GAT TCT GAG GGA ACT GTG AGC TTC CTC Leu Arg Pro Ile Val Val Lys Asp Ser Glu Gly Thr Val Ser Phe Leu 110

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							50							
		_		Lys				Leu					TTT	434
		GAA Glu 145	Gly				Arg					Val	AAG Lys	482
_		Ser				_				_	Lys		TTC Phe	530
	Ile	CAT His											CTT Leu	578
Сув		GAC Asp			Asn		 							62 6
_	_	GAA Glu						_						674
		TGG Trp 225												722
		AAC Asn												770
		CAA Gln												818
Ile		AAT Asn						_				_		866
		TCA Ser							_	_				914
		GCT Ala 305												962
		GAG Glu												1010
		GCT Ala												1058
		AAA Lys	Ile											1106
		ATT Ile												1154
	Phe	CAC His 385				Glu				Ser				1202

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			, Glu			GCA Ala					Tyr				1250
		Lys				GAA Glu 420									1298
	Lys					ACA Thr		 		 					1346
					Leu	CTT Leu									1394
			_	_	_	TGC Cys					_	_		;	1442
						TTC Phe							_		1490
						GGC Gly 500								;	1538
						ATG Met								:	1586
				_		GCA Ala			_					:	1634
						GTG Val								:	1682
			_			TTT Phe	_					_		:	1730
						ACT Thr 580								:	1778
						TGC Cys								:	1826
				_		TAT Tyr		-						:	1874
						GCT Ala								:	1922
CTG Leu						Leu								:	1970
GAT Asp					_									2	2018

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	Pro		GCC			Tyr				Met			20	66
			TGC Cys		Met								21	14
			TCC Ser 705	Ile								AAG Lys	21	62
			Leu									GGA Gly	22	10
			GAA Glu										22	158
			GAG Glu										23	106
			ACA Thr										23	54
			GAA Glu 785										24	02
			CCC Pro						_				24	50
			CAA Gln										24	98
			ACC Thr										25	46
			AAC Asn										25	94
		_	ATT Ile 865				_	-	_				26	36
TGAC	TCGA	.G											26	45

(2) INFORMATION FOR SEQ ID NO:23:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 875 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

Met Glu Arg Ala Gly Pro Ser Phe Gly Gln Gln Arg Gln Gln Gln Gln 15

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Pro Gln Gln Gln Lys Gln Gln Gln Arg Asp Gln Asp Ser Val Glu Ala Trp Leu Asp Asp His Trp Asp Phe Thr Phe Ser Tyr Phe Val Arg Lys Ala Thr Arg Glu Met Val Asn Ala Trp Phe Ala Glu Arg Val His Thr Ile Pro Val Cys Lys Glu Gly Ile Arg Gly His Thr Glu Ser Cys Ser 65 Cys Pro Leu Gln Gln Ser Pro Arg Ala Asp Asn Ser Val Pro Gly Thr Pro Thr Arg Lys Ile Ser Ala Ser Glu Phe Asp Arg Pro Leu Arg Pro 100 Ile Val Val Lys Asp Ser Glu Gly Thr Val Ser Phe Leu Ser Asp Ser 115 Glu Lys Lys Glu Gln Met Pro Leu Thr Pro Pro Arg Phe Asp His Asp 130 135 Glu Gly Asp Gln Cys Ser Arg Leu Leu Glu Leu Val Lys Asp Ile Ser 150 160 145 Ser His Leu Asp Val Thr Ala Leu Cys His Lys Ile Phe Leu His Ile 165 175 170 His Gly Leu Ile Ser Ala Asp Arg Tyr Ser Leu Phe Leu Val Cys Glu 180 Asp Ser Ser Asn Asp Lys Phe Leu Ile Ser Arg Leu Phe Asp Val Ala 195 Glu Gly Ser Thr Leu Glu Glu Val Ser Asn Asn Cys Ile Arg Leu Glu 210 215 220 Trp Asn Lys Gly Ile Val Gly His Val Ala Ala Leu Gly Glu Pro Leu 225 230 235 Asn Ile Lys Asp Ala Tyr Glu Asp Pro Arg Phe Asn Ala Glu Val Asp Gln Ile Thr Gly Tyr Lys Thr Gln Ser Ile Leu Cys Met Pro Ile Lys 265 Asn His Arg Glu Glu Val Val Gly Val Ala Gln Ala Ile Asn Lys Lys 275 Ser Gly Asn Gly Gly Thr Phe Thr Glu Lys Asp Glu Lys Asp Phe Ala 295 Ala Tyr Leu Ala Phe Cys Gly Ile Val Leu His Asn Ala Gln Leu Tyr 305 Glu Thr Ser Leu Leu Glu Asn Lys Arg Asn Gln Val Leu Leu Asp Leu 330 Ala Ser Leu Ile Phe Glu Glu Gln Gln Ser Leu Glu Val Ile Leu Lys Lys Ile Ala Ala Thr Ile Ile Ser Phe Met Gln Val Gln Lys Cys Thr

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Ile Phe Ile Val Asp Glu Asp Cys Ser Asp Ser Phe Ser Ser Val Phe . 370 His Met Glu Cys Glu Glu Leu Glu Lys Ser Ser Asp Thr Leu Thr Arg Glu His Asp Ala Asn Lys Ile Asn Tyr Met Tyr Ala Gln Tyr Val Lys Asn Thr Met Glu Pro Leu Asn Ile Pro Asp Val Ser Lys Asp Lys Arg 420 Phe Pro Trp Thr Thr Glu Asn Thr Gly Asn Val Asn Gln Gln Cys Ile 435 Arg Ser Leu Cys Thr Pro Ile Lys Asn Gly Lys Lys Asn Lys Val 450 455 Ile Gly Val Cys Gln Leu Val Asn Lys Met Glu Glu Asn Thr Gly Lys 465 Val Lys Pro Phe Asn Arg Asn Asp Glu Gln Phe Leu Glu Ala Phe Val Ile Phe Cys Gly Leu Gly Ile Gln Asn Thr Gln Met Tyr Glu Ala Val 505 Glu Arg Ala Met Ala Lys Gln Met Val Thr Leu Glu Val Leu Ser Tyr 520 His Ala Ser Ala Ala Glu Glu Glu Thr Arg Glu Leu Gln Ser Leu Ala 530 535 Ala Ala Val Val Pro Ser Ala Gln Thr Leu Lys Ile Thr Asp Phe Ser 545 550 Phe Ser Asp Phe Glu Leu Ser Asp Leu Glu Thr Ala Leu Cys Thr Ile 565 570 575 Arg Met Phe Thr Asp Leu Asn Leu Val Gln Asn Phe Gln Met Lys His 585 580 Glu Val Leu Cys Arg Trp Ile Leu Ser Val Lys Lys Asn Tyr Arg Lys Asn Val Ala Tyr His Asn Trp Arg His Ala Phe Asn Thr Ala Gln Cys 610 615 Met Phe Ala Ala Leu Lys Ala Gly Lys Ile Gln Asn Lys Leu Thr Asp 625 630 Leu Glu Ile Leu Ala Leu Leu Ile Ala Ala Leu Ser His Asp Leu Asp 645 His Arg Gly Val Asn Asn Ser Tyr Ile Gln Arg Ser Glu His Pro Leu 665 660 Ala Gln Leu Tyr Cys His Ser Ile Met Glu His His His Phe Asp Gln 685 675 Cys Leu Met Ile Leu Asn Ser Pro Gly Asn Gln Ile Leu Ser Gly Leu 690 695 Ser Ile Glu Glu Tyr Lys Thr Thr Leu Lys Ile Ile Lys Gln Ala Ile 710 715 720

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Leu Ala Thr Asp Leu Ala Leu Tyr Ile Lys Arg Arg Gly Glu Phe Phe Glu Leu Ile Arg Lys Asn Gln Phe Asn Leu Glu Asp Pro His Gln Lys Glu Leu Phe Leu Ala Met Leu Met Thr Ala Cys Asp Leu Ser Ala Ile Thr Lys Pro Trp Pro Ile Gln Gln Arg Ile Ala Glu Leu Val Ala Thr Glu Phe Phe Asp Gln Gly Asp Arg Glu Arg Lys Glu Leu Asn Ile Glu Pro Thr Asp Leu Met Asn Arg Glu Lys Lys Asn Lys Ile Pro Ser Met Gln Val Gly Phe Ile Asp Ala Ile Cys Leu Gln Leu Tyr Glu Ala Leu Thr His Val Ser Glu Asp Cys Phe Pro Leu Leu Asp Gly Cys Arg Lys Asn Arg Gln Lys Trp Gln Ala Leu Ala Glu Gln Gln Glu Lys Met Leu Ile Asn Gly Glu Ser Gly Gln Ala Lys Arg Asn

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Applicant's or agent's file reference number

32083

International application Notice 2017 US 9 4 / 06066

INDICATIONS RELATING TO A DEPOSITED MICROORGANISM

(PCT Rule 13bis)

A The indications and below the	formal to us the single self-self-self-self-self-self-self-self-
A. The indications made below relate to the microorganism re on page	•
B. IDENTIFICATION OF DEPOSIT	Further deposits are identified on an additional sheet
Name of depositary institution	
American Type Culture Collection	
Address of depositary institution (including postal code and country	vj
12301 Parklawn Drive Rockville, Maryland 20852 US	
Date of deposit	Accession Number
4 May 1993	ATCC 69296
C. ADDITIONAL INDICATIONS (leave blank if not applicab	ole) This information is continued on an additional sheet
a sample of the deposited microorganis publication of the mention of the gran date on which the application has been be withdrawn, only by the issue of sucthe person requesting the sample (Rule	t of the European patent or until the refused or withdrawn or is deemed to h a sample to an expert nominated by
D. DESIGNATED STATES FOR WHICH INDICATIO	NS ARE MADE (if the indications are not for all designated States)
EP	
E. SEPARATE FURNISHING OF INDICATIONS (leave	blank if not applicable)
The indications listed below will be submitted to the International lumber of Deposit*)	Bureau later (specify the general nature of the indications e.g., "Accession
For receiving Office use only	For International Bureau use only
This sheet was received with the international application	This sheet was received by the International Bureau on:
Doris L. Brock Division FOT INTERNATIONAL Division	Authorized officer

Form PCT/RO/134 (July 1992)

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CLAIMS

- 1. A purified and isolated polynucleotide encoding cGB-PDE.
- 2. The polynucleotide of claim 1 which is a DNA sequence.
- 3. The DNA sequence of claim 2 which is a cDNA sequence or a biological replica thereof.
- 4. The DNA sequence of claim 2 which is a genomic DNA sequence or a biological replica thereof.
 - 5. An RNA transcript of the genomic DNA sequence of claim 4.
- 6. The DNA sequence of claim 2 which is a wholly or partially chemically synthesized DNA sequence or a biological replica thereof.
- 7. The DNA sequence of claim 4 further comprising an endogenous expression control DNA sequence.
- 8. A DNA vector comprising a DNA sequence according to claim
- 9. The vector of claim 8 wherein said DNA sequence is operatively linked to an expression control DNA sequence.
- 10. A host cell stably transformed or transfected with a DNA sequence according to claim 7 in a manner allowing the expression in said host cell of cGB-PDE polypeptide possessing a ligand/receptor binding biological activity or immunological property specific to cGB-PDE.

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- 11. A method for producing cGB-PDE polypeptide, said method comprising growing a host cell according to claim 10 in a suitable nutrient medium and isolating cGB-PDE polypeptide from said cell or the medium of its growth.
- 12. A polypeptide or peptide capable of specifically binding to cGB-PDE.
 - 13. An antibody substance according to claim 12.
 - 14. A monoclonal antibody according to claim 13.
- 15. A hybridoma cell line producing a monoclonal antibody according to claim 14.
 - 16. A humanized antibody substance according to claim 13.
- 17. An antisense polynucleotide specific for a polynucleotide encoding cGB-PDE.
- 18. A DNA sequence encoding cGB-PDE and selected from the group consisting of:
 - (a) the DNA sequence set out in SEQ ID NO: 9 or 22;
- (b) a DNA which hybridizes under stringent conditions to the DNA of (a); and
- (c) a DNA sequence which, but for the redundancy of the genetic code, would hybridize under stringent conditions to a DNA sequence of (a) or (b).

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FIGURE

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582 580 580 580 125 48	674 628 709 174 97
AALKAGKIOK SLLVTGKLKR TLLMTGKLKS TLLMTGRLKK LLYKNLELTN YIMLHTGIMH VLLGTPALEA VLLGTPALEA VLLGTPALEG	GS. SILERH GS. SILERH GS. SILERH SSEGSVMERH NDRSVLENH NDRSVLENH NDESVLENH NDESVLENH - NDESVLENH - NDESVLENH - NDESVLENH
HAFNYGOTMF HGFNYGOTMF HGFNYGOTMF HAFSVSHFCY HAADVTOTVH HAADVTOTVH HAADVTOSTN HAADVTOSTN HAADVTOSTN HAADVTOSTN	RSEHPLAQLY KSONPLAKLH KSONPLAKLH KSTSPLARLH ASKSVLAALY OTRSDVAILY OTRSDVAILY NTNSELALMY NSSSELALMY
K. NVAYHNWR R. ITYHNWR A. VTYHNWR D. P. PYHNWR KYKNPYHNUI KYKNPYHNOI KYKNPYHNOI KYKNPYHNOI KYKNPYHNOI K. DNPFHNSI K. DNPFHNSI	HRGTNNLYOM HRGTNNLYOM HRGTNNLYOM HRGTNNSFOV HIGTTNNFHI HIGTTNSFUI HPGVSNOFLI HPGVSNOFLI HPGLTNOFLY
WILSVKKNYR FMYSLSKGYR FLFSVSKGYR WMTYVRKGYR FCLMVKKGYR FLDALETGYG YLLTLEGHYH FMSTLEDHYV	LIAALSHDLD VTAAFCHDID VTAGLCHDID LAAAFCHDID FISCMCHDLD VFAAAIHDVE IFACAIHDVD LFAACIHDVD
FOMKHEVLCK FUIPOEALVR FOIPOEVLVR FKYPVEVLTR YKIDCPTLAR FKIPYSCLIA FKIPTVFLMT FOIPADTLLR MIPPKTFLN	RLTDLEALAM YFTDLEAFAM YYTDLEAFAM YYTDLEAFAM YLEDMEIFAL WLTELEILAM CLSEIEVLAI VFTDLEVGGA VFTPLEVGGA
cGB-PDE ROS-a ROS-a CONE-a' cGS 61 KCAM 63 KCAM RATDUNCE DROSDUNCE CONSERVED	CGB-PDE ROS-B CONE-a' CGS 61 KCAM 63 KCAM RATDUNCE DROSDUNCE CONSERVED

FIGURE

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714 680 678 326 328 143	764 728 728 728 376 378 193
LATD LALYEKRETH IATD LALYEKRETH IATD LALYFKRETH LATD LAHHLRIFKD LATD MSGHFOOTKN LATD MSCHFOOVKS LATD MSKHMSLLAD LSTD MSKHMSLLAD ASTD MSKHMSLLAD LSTD MSKHMSLLAD LSTD MSKHMSLLAD	TACD LSAITKPWEY ITACD LSAITKPLPL ITACD LSNPAKPLPL ITACD L
LYIEMVIDAY MYIDAY MYIDAY MYIDAY MYIDAY MYIDAY	FLAMIN VMAMMI INAMMI THELL VLOSLV VLENLY
SIEEYKTTLK NRROHEHAIH NRROHEHVIH SRKDYORMLD SKDDWRDLRN TKDEFVELRA STKOKLSLRK OKKOROTLRK	NLEDPHOKEL MMLDOTRKEI LSLETTRKEI VTIDPTKKEI DRTNKOHHSL DRTNKOHHSL LLDNYSDRIO LLDNYSDRIO LLDNYTDRIO
SPGNOILSGL DESLNIFONL DESLNIFONL THGCNIFONL DESLNIFONL GENCOIFONL GENCOIFONL NGGCDIFONL	YETGGEWTOY YEDRKSWVEY METEEEAIKY L J SLGVL A GSGVL
HEDOCLMILN HLEFGKTLLR HLEYSKTLLO HLEYSKTLLO HVSAAYRLMO HISSVFRMO HLAVGFKLLO HLAVGFKLLO HLAVAFKLLO	FELIMKN FOKIVDOSKT FOKIVDESKN FOKIVDACEK LOKMAE IRNSLOOLER LKTMVETKKV LKTMVETKKV
cGB-PDE ROS-A ROS-B CONE-a' cGS GS KCAM 63 KCAM RATDUNCE DROSDUNCE CONSERVED	cGB-PDE ROS-a ROS-b CONE-a' cGS 61 KCAM 63 KCAM RATDUNCE DROSDUNCE CONSERVED

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812 778 778 776 844 422 424 339 3 / 10

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FIGURE

SMOVGFID KLOVGFID KLOVGFID KLOVGFID OSOIGFID OSOIGFID KSOVGFID MORNKADELP MDRNKADELP MDRNKKDELP MDREKAY. IP LCDRKSTMVA LCDRKSTMVA LCDRKSTMVA MCDRHTASVE MCDRHNATIE KELNIEPADL TVLOONPIPM TVLOOOPIPM A. MGNRPMEM EL. GLPFSP EL. GLPFSP ES. GLDISP ES. GNDISP OGDREA OGDLER OGDREA OGDLER OSKVALLVAA OSKVALLVAA OSQVALLVAN TRKIAELIYK HHRWTMALME HSRWTKALME YROWTERIMA YKRWVALLME

DROSDUNCE

ROS-a ROS-B CONE-a' cGS 61 KCAM 63 KCAM RATDUNCE

cGB-PDE

CONSERVED

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188 106 107 109	237 292 154 155	287 342 204 205 207
LELVS EDN. LO. LS MFLCR ARNGTPE. VA SLFMYR ORNGVAE. LA SLFMYR ARNGIAE. LA	/AAFGE PLNIKDAYED /VEDKK SIQLKDLTSE /AHTKK TFNVPDVKKN /AQTKK MVNVQDVMEC /ALSKK IVNVPNTEED	ACAFNKLGGDLFTD FMAVNKVDASEFSK IMAVNKLDGPCFTS IMAVNKLDGPCFTS IMAVNKVDGPHFTE
FLHIHGLIS ADRYS VLOYLOOETO ASRCC ALORLAOLLO ADRCS VMKKLCFLLO ADRMS VMKKLCFLLO ADRMS	NCIRLEWNKG IVGHV, EISFPLTTG RLGQV REAVFPLDVG IVGWV, SEIVFPLDIG VVGHV, SEIVFPLDMG VVGHV,	WPIKNHR.EE VVGV/VPVISRATDO VVAL/TPIV.MGKE VLAVISPIM.NGKD VVAVISPIM.NGKD VVAVI
HLDVTALCHK I LDASSLOLK VI AGSVELAAHR AI NVNMERVVFK I	TLEEASN I VLEEASN I KFEDNLVVPD I VLEECLVAPD I VLEECLVAPD I	TGYKTOSILC LGCEVOAMLC TGYVTRNLLA TDYVTRNILA TEYOTKNILA
LLELVKDISS ILOLCGELYD LLEVLLEE LFELVODMOE LLRDFOD	SRLFDVAEGS CKVIGDK SKLLDVTPTS TRLFSVQPDS TRLFNVHKDA	PRFNAEVDOI DM. QOLOSM SHFSDFMDKO PHFSSFADEL EHFCDFVDTL
cGB-PDE cGS CONE-a' ROS-B ROS-a CONSERVED	cGB-PDE cGS CONE-a' ROS-B ROS-a CONSERVED	cGB-PDE cGS CONE-a' ROS-β ROS-α CONSERVED

cGB-PDE cGS CONE-a' ROS-B ROS-a CONSERVED	KDEKDFAAYL ODEHVIOHCF ODEEVFSKYL EDEDVFLKYL NDEEILLKYL -DE	AFCGIVLHNA HYTSTVLTST SFVSIILKLH NFGTLNLKIY NFGTLNLKIY	OLYETSLLEN LAFOKEOKLK HTNYLYNIES HYSYLHNCET HLSYLHNCET	KRNOVLLDLA CECOALLOVA RRSOILMWSA RRGOVLLWSA RRGOILLWSG	SLIFEEQOSL KNLFTHLDDV NKVFEELTDV NKVFEELTDI SKVFEELTDI	337 390 252 253 255	WO 94/28144
-PDE E-α' E-α' SERVED	EVILKKIAAT SVLLOEIITE EROFHKALYT EROFHKAFYT EROFHKALYT	IISFMOVOKC ARNLSNAEIC VRTYLNCERY VRAYLNCDRY VRAFLNCDRY	TIFIVD. EDC SVFLID 0 SIGLLDMTKE SVGLLDMTKE SVGLLDMTKQ	SDSFSSVFHM NELVAKVFDG KEFY. DEWPV KEFF. DVWPV KEFF. DVWPV	ECEELEKSSD GVLEDESY KPGEVEPYKG LMGEAPPYAG LMGEAPPYAG	361 301 302 304	
cGB-PDE cGS CONE-a' ROS-B ROS-a CONSERVED	TLTRE PKTPDGREVI PRTPDGREIL PRTPDGREIL	FYKIIDYILH FYKVIDYILH FYKVIDYILH	GKEEIKVIPT GKEDIKVIPS GKEDIKVIPS	PADOPADO	MYAQYVKNTM GIAGHVATTG GLPTYVAENG GLPTYVAESG GLPTYVAQNG	411 351 352 354	5 / 1 0

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FIGURE 2C

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459 499 400 402	506 541 441 442	526 561 461 462 464
KKNKVIGVCO ENQEVIGVAE KKEDIVGVAT KKEEIVGVAT KKEEIVGVAT	EAVERAMAKO KKVNEAQYRS EKMNKLENRK DKMNKLENRK ELMNKLENRK	
SLLCTPIKNG NILCFPIKN. NVLSLPIVN. NVLSMPIVN. NVLSMPIVN. LPI-N-	GLGIONTOMY GISIAHSLLY GWSLLNTDTY GWSVLNTDTY GWSVLNTDTY GWSVLNPDTY GY	
MGNINOOCIR DDSTGRF. TR VDETGWV. IK LDDSGWI. VK LDESGWM. IK	OFLEAFVIFC DLATAFSIYC HIAETLTOFL VLMESLTOFL TLMESLAOFL	
DKRFPWTNEN HPLFYRGV DEYFTFOKGP DEMFNFOEGP EDFFAFOKEP	KVKAFNRNDE PWFSKFDE KPFDEYDE KPFDEODE KPFDEMDE	ASAAEEE MKVSDDE TKATPDE VRCDREE VKCDNEE
EPLNIPDVSK OILNIPDAYA FICNMLNAPA FICNIMNAPA LICNIMNAPS LICNIMNAPS	LVNKMEETTG LVNKING FYNRKDG FYNRKDG	MVTLEVLSYH HLANEMMMYH DIAQEMLMNH DIAQDMVLYH DIFQDMVKYH
cGB-PDE cGS CONE-a' ROS-B ROS-A CONSERVED	cGB-PDE cGS CONE-a' ROS-B ROS-a CONSERVED	cGB-PDE cGS CONE-a' ROS-B ROS-a Conserved

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FIGURE 3

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EPLNIKDAYEDPRFNAEVDQITGYKTQSILCMPIKMH.REEVVGVAQAIN.KKSGN KIVNVPNTEEDEHFCDFVDTLTEYQTKNILASPIMNG.K.DVVAIIMAVN.KVDGP KIVNVPNTEEDEHFSSFADELTDYVTRNILATPIMNG.K.DVVAIIMAVN.KLDGP KAVNVQDVMECPHFSDFMDKQTGYVTRNILATPIMNG.K.DVVAVIMAVN.KLDGP KSIQLKDLTSEDMQQLQSMLGCEVQAMLCVPVISRATDQVVALACAFN.KLGGD EPLNIPDVSKDKRFPWTNENMGNINQQCIRSLLCTPIKNGKKNKVIGVCQLVN.KMEET LICNIMNAPSEDFFAFQKEPLDE.SGWMIKNVLSMPIVNK.KEEIVGVATFYNRKDGKP FICNIMNAPSEDFFAFQKEPLDD.SGWIVKNVLSMPIVNK.KEEIVGVATFYNRKDGKP FICNIMNAPADEYFTFQKGPVDE.TGWVIKNVLSMPIVNK.KEDIVGVATFYNRKDGKP QILNIPDAYAHPLFYRGVDDSTGFRTRNILCFPIKNE.NQEVIGVAELVN.KINGP	GGTFTEKDEKDFAAYLAFCGIVLHMAQL.YEHFTENDEEILLKYLNFANLIMKVFHLSYCFTSEDEDVFLKYLNFGTLNLKIYHLSYEFSKQDEEVFSKYLSFVSIILKLHHTNYLFTDQDEHVIQHCFHYTSTVL.TSTLAFQ TGKVKAFNRNDEQFLEAFVIFCGLGIQNTQM.YEFVEQDEVLMESLTQFLGWSV.LNTDTYDFVEQDEVLMESLTQFLGWSV.LNTDTYEFVEQDEVLATAFSIYCGISI.AHSLLYKFDEYDEHIAETLTQFLGWSL.LNTDTYE
CGB-PDE ROS-a ROS-a CGB-PDE CGB-PDE ROS-a ROS-a CGS CGS CGS CGS CGS CGS CGS CGS CGS	CGB-PDE ROS-a CGS-a CGB-PDE CGS-a CGS-a CGS-a CGS-a CGS-a

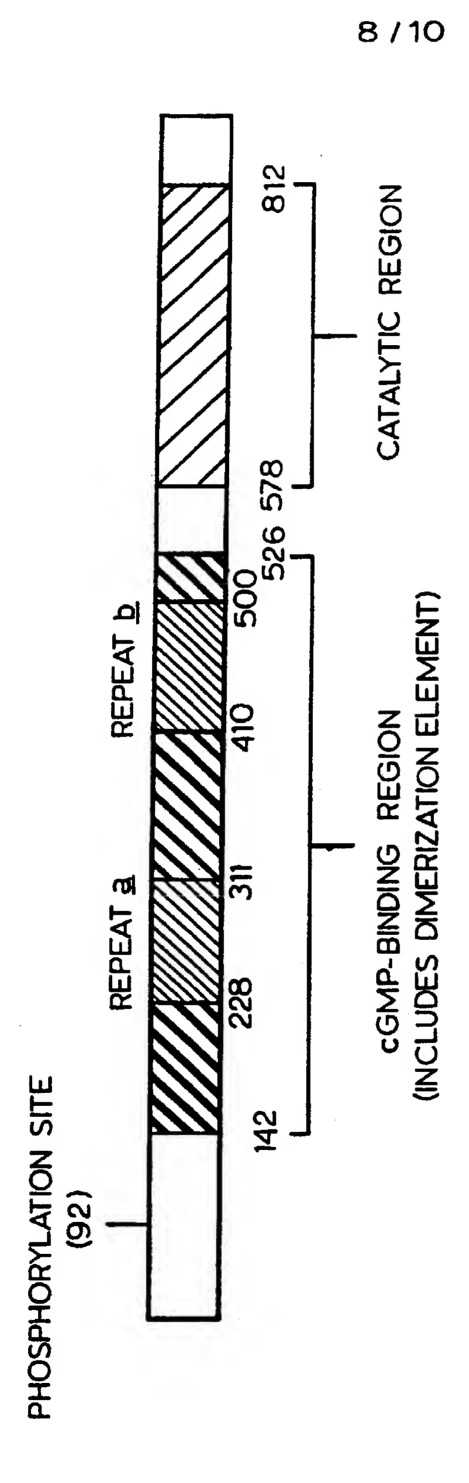
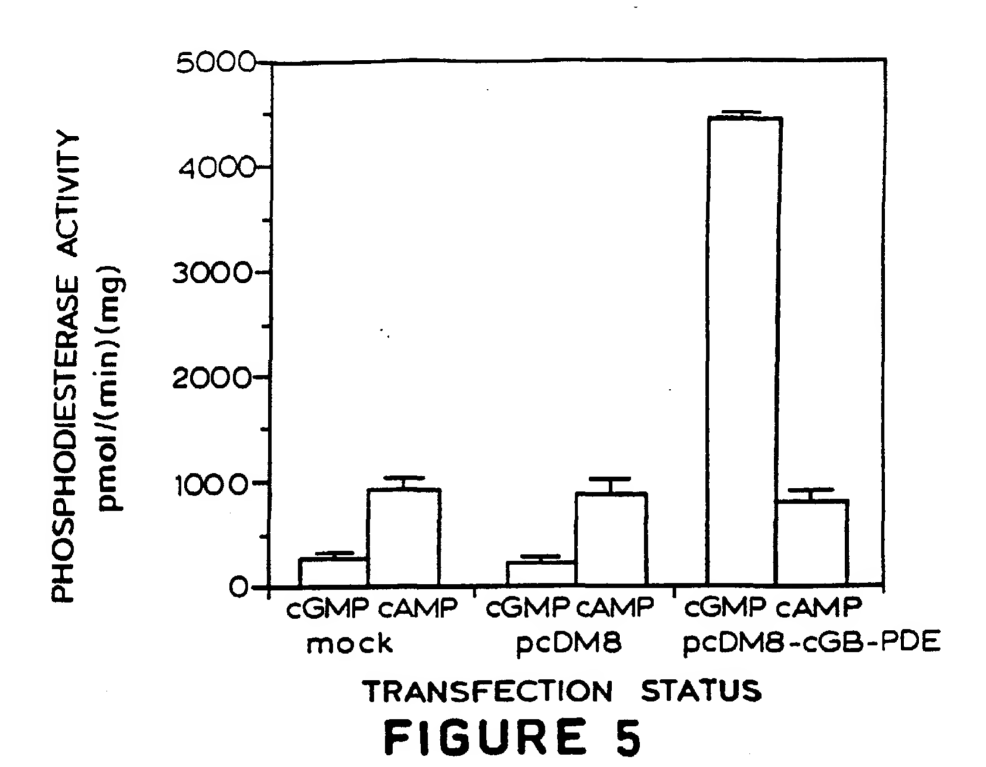
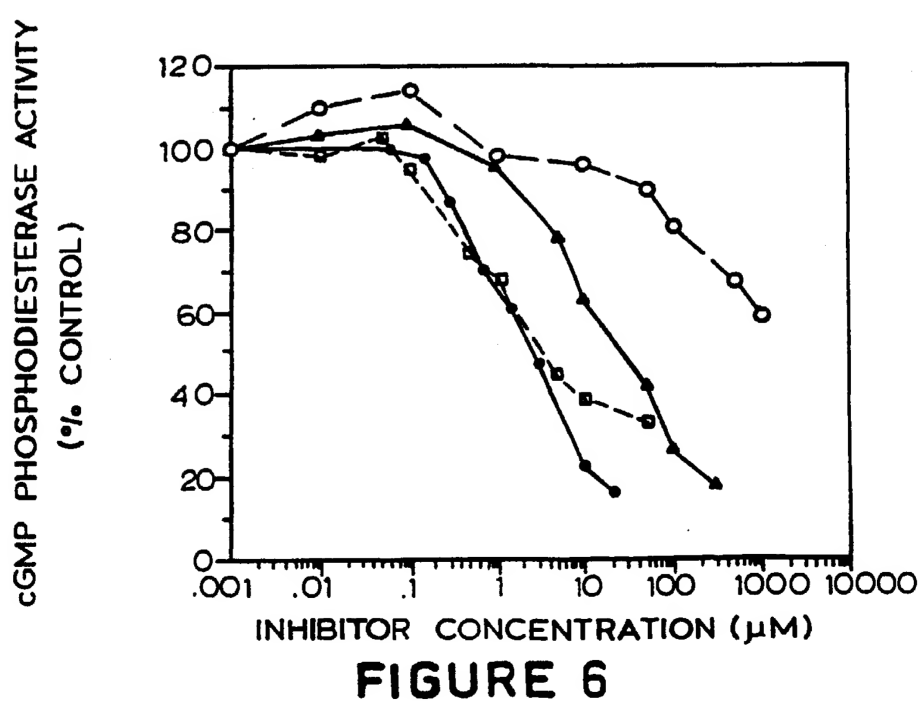


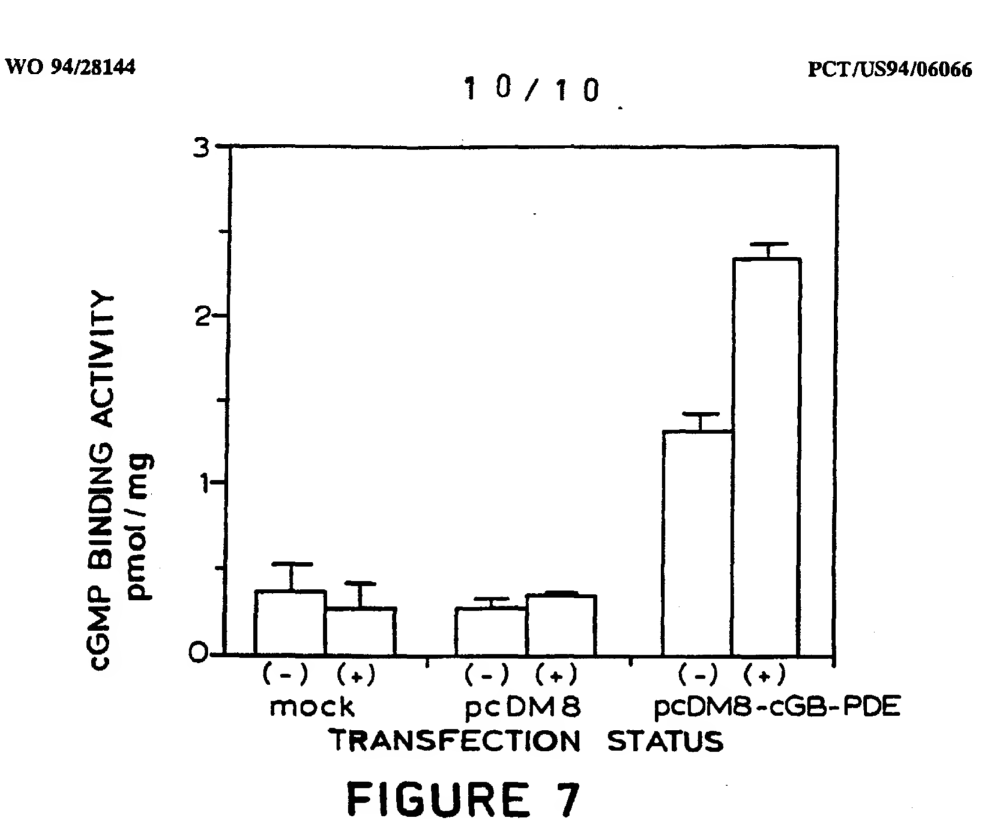
FIGURE 4

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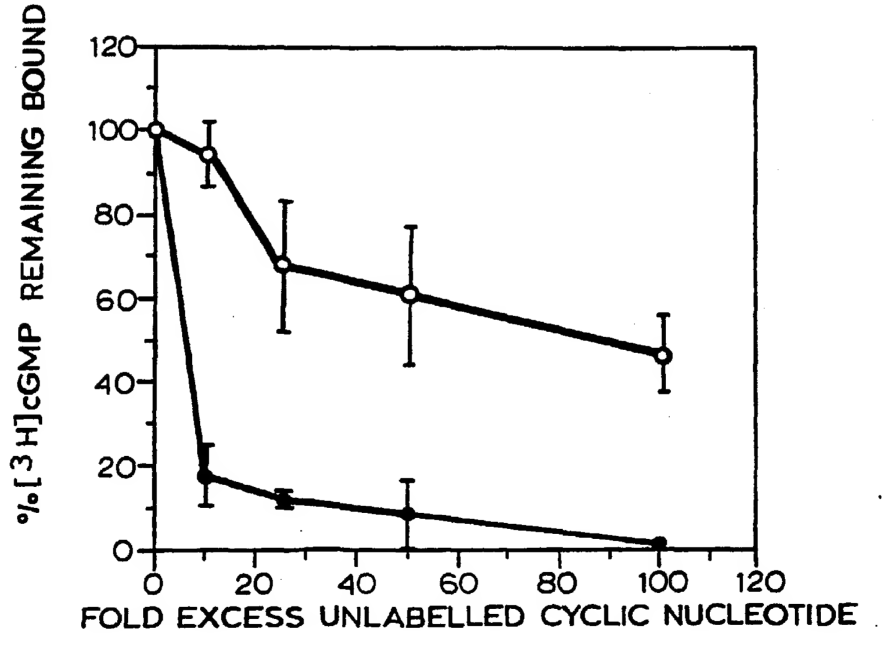


FIGURE 8

	INTERNATIONAL SEARCH R	REPORT	Int and Appl PCT/US 94	lication No
A. CL.'S	SIFICATION OF SUBJECT MATTER			
IPC 5	C12N15/55 C12N9/16 C12P21/	08 C12N15	/11	
According	to International Patent Classification (IPC) or to both national class	ification and IPC		
	S SEARCHED			
Minimum IPC 5	documentation searched (classification system followed by classification s	tion symbols)		
Documenta	ation searched other than minimum documentation to the extent that	such documents are in	cluded in the fields a	earched
Electronic	data hase consulted during the international search (name of data ha	se and, where practica	i, search terms used)	
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C, DOCUM	MENTS CONSIDERED TO BE RELEVANT	······································		
Category *	Citation of document, with indication, where appropriate, of the	elevant passages		Relevant to claim No.
Υ	JOURNAL OF BIOLOGICAL CHEMISTRY.	-		1-18
	vol.265, no.25, 5 September 1990 BALTIMORE US	•		
	pages 14971 - 14978			
	THOMAS ETAL. 'Substrate- and			
	Kinase-directed regulation of Phosphorylation of a cGMP-binding	~		
	Phosphodiesterase by cGMP.	9		
	cited in the application			
	see the whole document			
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X Furt	her documents are listed in the continuation of box C.	Patent family	members are listed	in annex.
* Special cat	tegories of cited documents:	T later document pr		
'A' docum	ent defining the general state of the art which is not ered to be of particular relevance	cited to understan	ind not in conflict wi nd the principle or th	th the application but seory underlying the
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filing of the L' docume	ent which may throw doubts on priority claim(s) or	involve an inven	ered novel or cannot tive step when the do	cument is taken alone
citation	is cited to establish the publication date of another n or other special reason (as specified)	cannot be consid	icular relevance; the ered to involve an in	ventive step when the
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"P" docume	ent published prior to the international filing date but an the priority date claimed	in the art. "&" document membe	er of the same patent	family
Date of the	actual completion of the international search	Date of mailing o	f the international se	arch report
20	6 September 1994		1 9, 10.	94
Name and n	nailing address of the ISA	Authorized office		
	European Patent Office, P.B. 581 8 Patentiaan 2 NL - 2280 HV Rijswijk	_		
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INTERNATIONAL SEARCH REPORT

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C(Continu	ation) DOCUMENTS CONSIDERED TO BE RELEVANT	PCT/US 94/06066	
itegory *		Relevant to claim No.	
	JOURNAL OF BIOLOGICAL CHEMISTRY., vol.265, no.25, 5 September 1990, BALTIMORE, MD US pages 14964 - 14970 THOMAS ET AL. 'Characterization of a Purified Bovine Lung cGMP-binding cGMP Phosphodiesterase' cited in the application see the whole document	1-18	
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	GENOMICS, vol.13, no.3, July 1992 pages 698 - 704 COLLINS ET AL. 'The human Beta-Subunit of Rod Photoreceptor cGMP Phosphodiesterase: Complete Retinal cDNA Sequence and Evidence for Expression in Brain' cited in the application see the whole document	1-18	
, X	JOURNAL OF BIOLOGICAL CHEMISTRY., vol.268, no.30, 25 October 1993, BALTIMORE US pages 22863 - 22873 MCALLISTER-LUCAS ET AL. 'The Structure of a Bovine Lung cGMP-binding, cGMP-specific Phosphodiesterase Deduced from a cDNA Clone' see the whole document	1-18	